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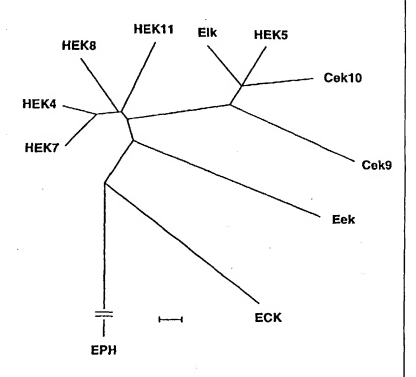
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(54) Title: HEK5, HEK7, HEK8, HEK11, NEW EPH-LIKE RECEPTOR PROTEIN TYROSINE KINASES

(57) Abstract

Four novel members of the EPH subfamily of receptor protein tyrosine kinases are disclosed. Nucleic acid sequences encoding receptor proteins, recombinant plasmids and host cells for expression, and methods of producing and using such receptors are also disclosed.



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HEK5, HEK7, HEK8, HEK11, new EPH-like receptor protein tyrosine kinases

Field of the Invention

5 The invention relates generally to receptor protein tyrosine kinases (PTKs) and particularly to novel Eph-like receptor PTKs, to fragments and analogs thereof, and to nucleic acids encoding same. The present invention also relates to methods of producing and using such receptors.

Background of the Invention

Receptor PTKs are a structurally related family of proteins that mediate the response of cells to 15 extracellular signals (Ullrich et al. Cell 61, 203-212 These receptors are characterized by three major functional domains: an intracellular region containing the sequences responsible for catalytic activity, a single hydrophobic membrane-spanning domain, 20 and a glycosylated extracellular region whose structure determines ligand binding specificity. Signal transduction is initiated by the binding of growth or differentiation factors to the extracellular domain of their cognate receptors. Ligand binding facilitates 25 dimerization of the receptor which can induce receptor autophosphorylation. Both soluble and membraneassociated protein ligands have been shown to function in this manner. This process is the initial step in a cascade of interactions involving the phosphorylation of 30 a variety of cytoplasmic substrates and culminating in a biological response by the cell. The best characterized response to tyrosine kinase receptor activation is cell growth. However, analysis of the role of some growth factors in vivo suggests that differentiation or cell 35

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survival might also be mediated by tyrosine kinase receptor/ligand interactions.

Receptor PTKs have been grouped into fairly well-defined families on the basis of both sequence homology and shared structural motifs. The amino acid sequence of the portion of the intracellular domain responsible for the catalytic activity is well conserved among all tyrosine kinases and even more closely matched within a receptor sub-family. Comparisons of this 10 portion of the amino acid sequence have been used to construct phylogenetic trees depicting the relatedness of family members to each other and to the tyrosine kinases as a whole (Hanks and Quinn, Methods Enzymol. 200, 38-62 (1991)). This sequence conservation has also 15 been exploited in order to isolate new tyrosine kinases using the polymerase chain reaction (PCR) (Wilks, Proc. Natl. Acad. Sci. USA 86, 1603-1607 (1989)). Oligonucleotides based on the highly conserved catalytic 20 domain of PTKs can be used as PCR primers to amplify related sequences present in the template. These fragments can then be used as probes for isolation of the corresponding full-length receptor clones from cDNA libraries. Anti-phosphotyrosine antibodies have also been used to identify PTK cDNA clones in phage 25 expression libraries (Lindberg and Pasquale, Methods Enzymol. 200, 557-564 (1991)). These strategies have been used by a number of investigators to identify an ever-increasing number of protein tyrosine kinase 30 receptors.

There are now 51 distinct PTK receptor genes that have been published and divided into 14 sub-families One such sub-family is the EPH-like receptors. The prototype member, EPH, was isolated by Hirai et.al. (Science 238, 1717-1720 (1987)) using low

stringency hybridization to a probe derived from the viral oncogene v-fps. EPH-like receptors have been implicated in cell growth based in part on studies which show that overexpression of the gene in NIH3T3 cells causes focus formation in soft agar and tumors in nude mice (Maru et al. Oncogene 5, 199-204 (1990)). Other members of the EPH sub-family which have been identified include the following:

ECK (Lindberg et al. Mol. Cell. Biol. 10,

10 6316-6324 (1990))

Elk (Lhoták et al. Mol. Cell. Biol. <u>11</u>, 2496-2502 (1991))

Ceks 4,5,6,7,8,9, and 10 (Pasquale, Cell Regulation $\underline{2}$, 523-534 (1991); Sajjadi et al. The New Biologist $\underline{3}$, 769-778 (1991); Sajjadi and Pasquale Oncogene $\underline{8}$, 1807-1813 (1993))

HEK2 (Bohme et al. Oncogene 8, 2857-2862 (1993))

Eek, Erk (Chan and Watt, Oncogene 6, 1057-1061

20 (1991))

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Ehk1, Ehk2 (Maisonpierre et al. Oncogene 8, 3277-3288 (1993))

Homologs for some of these receptors have been identified in other species (Wicks et al. Proc. Natl. 25 Acad. Sci. USA 89, 1611-1615 (1992)); Gilardi-Hebenstreit et al. Oncogene 7, 2499-2506 (1992)). expression patterns and developmental profiles of several family members suggest that these receptors and their ligands are important for the proliferation, 30 differentiation and maintenance of a variety of tissues (Nieto et al. Development 116, 1137-1150 (1992)). Structurally, EPH sub-family members are characterized by an Ig-like loop, a cysteine rich region, and two fibronectin-type repeats in their extracellular domains. 35 The amino acid sequences of the catalytic domains are

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more closely related to the SRC sub-family of cytoplasmic PTKs than to any of the receptor PTKs.

Among the catalytic domains of receptor PTKs, the EPH sub-family is most similar in amino acid sequence to the epidermal growth factor receptor sub-family.

It is an object of the invention to identify novel receptors belonging to the EPH sub-family. A directed PCR approach has been used to identify five human EPH-like receptors from a human fetal brain cDNA library. These receptors are designated HEK4, HEK5, HEK7, HEK8, and HEK11. The relationship of these receptors to previously identified EPH-like receptors is as follows:

HEK4 is the human homolog of Cek4 (chicken) and Mek4 (mouse) and is identical to HEK (Boyd et al. J. Biol. Chem. <u>267</u>, 3262-3267 (1992); Wicks et al., 1992) which was previously isolated from a human lymphoid tumor cell line.

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HEK5 is the human homolog of Cek5, a fulllength eph-like receptor clone from chicken. A portion of the HEK5 sequence was previously disclosed as ERK, a human clone encoding about sixty amino acids (Chan and Watt, 1991)

HEK7 is the human homolog of Cek7 isolated from chicken.

HEK8 is the human homolog of Cek8 a fulllength clone from chicken and Sek, a full-length clone from mouse. (Nieto et al., 1992; Sajjadi et al., 1991)

HEK11 does not have a known non-human homolog. With the addition of the new members HEK5, HEK7, HEK8 and HEK11 and the report of a PCR fragment encoding an eph-like receptor (Lai & Lemke Neuron 6, 691-704 (1991)), a total of twelve distinct sequences that represent EPH-like receptors have been published, making it the largest known sub-family of PTKs.

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It is a further object of the invention to generate soluble EPH-like receptors and antibodies to EPH-like receptors. Soluble receptors and antibodies are useful for modulating EPH-like receptor activation.

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Summary of the Invention

The present invention provides novel EPH-like receptor protein tyrosine kinases. More particularly, the invention provides isolated nucleic acids encoding four novel members of the sub-family of EPH-like receptor PTKs which are referred to collectively as HEKs (human-eph like kinases). Also encompassed are nucleic acids which hybridize under stringent conditions to EPH-like receptor nucleic acids. Expression vectors and host cells for the production of receptor polypeptides and methods of producing receptors are also provided.

Isolated polypeptides having amino acid sequences of EPH-like receptors are also provided, as are fragments and analogs thereof. Antibodies specifically binding the polypeptides of the invention are included. Also comprehended by the invention are methods of modulating the endogenous activity of an EPH-like receptor and methods for identifying receptor ligands.

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Description of the Figures

Figure 1 shows the nucleotide and predicted amino acid sequence of the HEK5 receptor.

30 Figure 2 shows the nucleotide and predicted amino acid sequence of the HEK7 receptor.

Figure 3 shows the nucleotide and predicted amino acid sequence of the HEK8 receptor.

Figure 4 shows the nucleotide and predicted amino acid sequence of the HEK11 receptor.

Figure 5 shows the comparison of the amino acid sequences of the human EPH receptor sub-family. multiple sequence alignment was done using the LineUp program included in the Genetics Computer Group sequence analysis software package (Genetics Computer Group, (1991), Program Manual for the GCG Package, Version 7, April 1991, Madison, Wisconsin, USA 53711). 10 indicate spaces introduced in order to optimize alignment. The predicted transmembrane domains and signal sequences of each receptor are indicated by underlining and italics, respectively. Cysteine 15 residues conserved throughout the sub-family are indicated with asterisks. Arrows indicate the tyrosine kinase catalytic domain. Amino acid sequences of EPH, ECK and HEK2 were taken from the appropriate literature references.

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Figure 6 shows the molecular phylogeny of the EPH subfamily of receptor protein tyrosine kinases. Catalytic domain sequences were analyzed as described by Hanks and Quinn, 1991. The scale bar represents an arbitrary evolutionary difference unit. The EPH branch, which has been shown with a discontinuity for the sake of compactness, is 23.5 units in length.

Figures 7-11 show Northern blot analyses of the tissue distribution of the HEK receptors. Receptor cDNA probes, labeled with ³²P, were hybridized to either 2 µg of poly A⁺ RNA from human tissues (panel A, Clontech) or 10 µg of total RNA from rat tissues (panel B). Sizes of the transcripts were determined by comparison with RNA molecular weight markers (Bethesda Research Labs,

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Gaithersburg, MD). Figure 7, HEK4; Figure 8, HEK5; Figure 9, HEK7; Figure 10; HEK8; Figure 11; HEK 11.

Detailed Description of the Invention

The present invention relates to novel 5 EPH-like receptor protein tyrosine kinases. More particularly, the invention relates to isolated nucleic acids encoding four novel members of the sub-family of EPH-like receptor PTKs. These four members are designated herein as HEK (human eph-like kinases). 10 Nucleic acids encoding HEK receptors were identified in a human fetal brain cDNA library using oligonucleotide probes to conserved regions of receptor PTKs and EPHlike receptor PTKs. The predicted amino acid sequences of three HEK receptors had extensive homology in the 15 catalytic domain to previously identified EPH-like receptors Cek5, Cek7 and Cek8 isolated from chicken and, accordingly, are designated HEK5, HEK7 and HEK8. predicted amino acid sequence of the fourth HEK receptor 20 revealed that it was not a homolog of any previously identified EPH-like receptor. It is designated HEK11. It is understood that the term "HEKs" comprises HEK5, HEK7, HEK8 and HEK11 as well as analogs, variants, and mutants thereof which fall within the scope of the 25 invention.

The invention encompasses isolated nucleic acids selected from the group consisting of:

- (a) the nucleic acids set forth in any of SEQ 30 ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14, or SEQ ID NO: 16 and their complementary strands;
 - (b) a nucleic acid hybridizing to the coding regions of the nucleic acids in any of SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14, or SEQ ID NO: 16 under stringent conditions; and

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(c) a nucleic acid of (b) which, but for the degeneracy of the genetic code, would hybridize to the coding regions of the nucleic acids in any of SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14, or SEQ ID NO: 16.

The nucleic acids of the invention preferably hybridize to HEK5, HEK7, HEK8, or HEK11 coding regions under conditions allowing up to about 5% nucleotide mismatch based upon observed nucleic acid identities among known human or nonhuman EPH-like receptors. An example of such a condition is hybridization at 60° in 1M Na+ followed by washing at 60° in 0.2XSSC. Other hybridization conditions may be ascertained by one skilled in the art which allow base pairing with similar levels of mismatch.

15 In a preferred embodiment, the isolated nucleic acids encode polypeptides having the amino acid sequences of HEK5, HEK7, HEK8 or HEK11. A nucleic acid includes cDNA, genomic DNA, synthetic DNA or RNA. Nucleic acids of this invention may encode full-length 20 receptor polypeptides having an extracellular ligand-binding domain, a transmembrane domain, and a cytoplasmic domain, or may encode fragments such as extracellular domains which are produced in a soluble, secreted form. Nucleic acid constructs which produce soluble HEK receptors are described in Example 3. 25 Polypeptides and fragments encoded by the nucleic acids have at least one of the biological activities of an EPH-like receptor protein tyrosine kinase, such as the ability to bind ligand.

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The invention also encompasses nucleic acids encoding chimeric proteins wherein said proteins comprise part of the amino acid sequence of a HEK receptor linked to an amino acid sequence from a heterologous protein. One example of such a chimeric protein is an extracellular domain of a HEK receptor

fused to a heterologous receptor cytoplasmic domain. Example 5 describes the construction and expression of a chimeric receptor comprising the HEK8 extracellular domain with the trkB cytoplasmic domain and a second chimeric receptor comprising the HEK11 extracellular domain with the trkB cytoplasmic domain. HEK receptors may also be fused to other functional protein domains, such as an Ig domain which acts as an antibody recognition site.

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The nucleic acids of the present invention may be linked to heterologous nucleic acids which provide expression of receptor PTKs. Such heterologous nucleic acids include biologically functional plasmids or viral vectors which provide genetic elements for 15 transcription, translation, amplification, secretion, One example of an expression vector suitable for producing EPH-like receptors of the present invention is $pDSR\alpha$ which is described in Example 3. It is understood 20 that other vectors are also suitable for expression of EPH-like receptors in mammalian, yeast, insect or bacterial cells. In addition, in vivo expression of nucleic acids encoding EPH-like receptor PTKs is also encompassed. For example, tissue-specific expression of EPH-like receptors in transgenic animals may be readily 25 effected using vectors which are functional in selected tissues.

Host cells for the expression of EPH-like
receptor PTKs will preferably be established mammalian cell lines, such as Chinese Hamster Ovary (CHO) cells or NIH 3T3 cells, although other cell lines suitable for expression of mammalian genes are readily available and may also be used. Such host cells are transformed or transfected with nucleic acid constructs suitable for expression of an EPH-like receptor. Transformed or

transfected host cells may be used to produce suitable quantities of receptor for diagnostic or therapeutic uses and to effect targeted expression of EPH-like receptors in selected adult tissues, such as brain, kidney, and liver, or in embryonic or rapidly dividing tissues.

The present invention provides purified and isolated polypeptides having at least one of the biological properties of an EPH-like receptor (e.g. 10 ligand binding, signal transduction). The isolated polypeptides will preferably have an amino acid sequence as shown in any of SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14 or SEQ ID NO: 16. Polypeptides of this invention 15 may be full-length polypeptides having an extracellular domain, a transmembrane domain, and a cytoplasmic domain, or may be fragments thereof, e.g., those having only an extracellular domain or a portion thereof. will be understood that the receptor polypeptides may 20 also be analogs or naturally-occurring variants of the amino acid sequences shown in SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14 or SEQ ID NO: 16. Such analogs are generated by amino acid substitutions, deletions and/or insertions using methods available in the art.

Polypeptides of the invention are preferably the product of expression of an exogenous DNA sequences, i.e., EPH-like receptors are preferably produced by recombinant means. Methods of producing EPH-like receptors comprising culturing host cells which have been transformed or transfected with vectors expressing an EPH-like receptor are also encompassed. EPH-like receptors, particularly fragments, may also be produced by chemical synthesis. The polypeptides so produced may be glycosylated or nonglycosylated depending upon the host cell employed, or may have a methionine residue at the amino terminal end. The polypeptides so produced

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are identified and recovered from cell cultures employing methods which are conventional in the art.

EPH-like receptors of the present invention are used for the production of antibodies to the receptors. Antibodies to HEK receptors have been described in Example 4. Antibodies which recognize the polypeptides of the invention may be polyclonal or monoclonal and may be binding fragments or chimeric antibodies. Such antibodies are useful in the detection of EPH-like receptors in diagnostic assays in the purification of receptor, and in the modulation of EPH-like receptor activation.

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As described in co-pending and co-owned U.S. Serial No. 08/145,616, the only known ligand for an 15 EPH-like receptor is a protein which binds to and induces phosphorylation of the eck receptor. receptor ligand was previously identified as B61. (Holzman et al. Mol. Cell. Biol. 10, 5830-5838 (1990)). The availability of ECK receptor was important for the 20 identification of a ligand since B61, although known, had not been previously implicated as an ECK receptor Therefore, EPH-like receptors having ligand binding domains are useful for the identification and purification of ligands. Polypeptides of the present 25 invention may be used to identify and purify ligands for HEK5, HEK7, HEK8 and HEK11 receptors. Binding assays for the detection of potential ligands may be carried out in solution or by receptor immobilization on a solid support using methods such as those described in 30 co-pending and co-owned U.S. Serial No. 08/145,616. Such assays may employ an isolated ligand binding domain of a HEK receptor. Alternatively, a HEK ligand binding domain fused to an Ig domain may be used to detect the presence of HEK ligand on cell surfaces. 35

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Soluble EPH-like receptors may be used to modulate (i.e., increase or decrease) the activation of the cell-associated receptors, typically by competing with the receptor for unbound ligand. Modulation of EPH-like receptor activation may in turn alter the proliferation and/or differentiation of receptor-bearing cells. For example, based upon the observed tissue distribution of the receptors of this invention (see Table 5), soluble HEK7 receptor is likely to primarily affect proliferation and/or differentiation of brain cells, while soluble HEK5 receptor may affect primarily brain and pancreatic cells, although effects of HEK5 receptor on other tissues may not be excluded.

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Antibodies to EPH-like receptors are useful 15 reagents for the detection of receptors in different cell types using immunoassays conventional to the art. Antibodies are also useful therapeutic agents for modulating receptor activation. Antibodies may bind to the receptor so as to directly or indirectly block 20 ligand binding and thereby act as an antagonist of receptor activation. Alternatively, antibodies may act as an agonist by binding to receptor so as to faciliate ligand binding and bring about receptor activation at lower ligand concentrations. In addition, antibodies of 25 the present invention may themselves act as a ligands by inducing receptor activation. It is also contemplated that antibodies to EPH-like receptors are useful for selection of cell populations enriched for EPH-like receptor bearing cells. Such populations may be useful in cellular therapy regimens where it is necessary to 30 treat patients which are depleted for certain cell types.

The isolated nucleic acids of the present inventions may be used in hybridization assays for the detection and quantitation of DNA and/or RNA coding for HEK5, HEK7, HEK8, HEK11 and related receptors. Such

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assays are important in determining the potential of various cell types to express these receptors and in determining actual expression levels of HEK receptors. In addition, the nucleic acids are useful for detecting abnormalities in HEK receptor genes, such as translocations, rearrangements, duplications, etc.

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Therapeutic regimens involving EPH-like receptors will typically involve use of the soluble form of the receptor contained in a pharmaceutical 10 composition. Such pharmaecutical compositions may contain pharmaceutically acceptable carrier, diluents, fillers, salts, buffers, stabilizers and/or other materials well known in the art. Further examples of such constituents are described in Remington's 15 Pharmaceutical Sciences 18th ed., A.R. Gennaro, ed. (1990). Administration of soluble EPH-like receptor compositions may be by a variety of routes depending upon the condition being treated, although typically administration will occur by intravenous or subcutaneous 20 methods. Pharmaceutical compositions containing antibodies to EPH-like receptors will preferably include mouse-human chimeric antibodies or CDR-grafted antibodies in order to minimize the potential for an · 25 immune response by the patient to antibodies raised in mice. Other components of anti-EPH antibody compositions will be similar to those described for soluble receptor.

The amount of soluble Eph-like receptors or anti-Eph antibody in a pharmaceutical composition will depend upon the nature and severity of the condition being treated. Said amount may be determined for a given patient by one skilled in the art. It is contemplated that the pharmaceutical compositions of the present invention will contain about 0.01 µg to about

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100 mg of soluble receptor or anti-Eph antibody per kg body weight.

A method for modulating the activation of an EPH-like receptor PTK is also provided by the invention. 5 In practicing this method, a therapeutically effective amount of a soluble EPH-like receptor or an anti-EPH antibody is administered. The term "therapeutically effective amount" is that amount which effects an increase or decrease in the activation of an EPH-like 10 receptor and will range from about 0.01 μ g to about 100 mg of soluble receptor or anti-EPH antibody per kg body weight. In general, therapy will be appropriate for a patient having a condition treatable by soluble receptor or anti-EPH antibody and it is contemplated that such a 15 condition will in part be related to the state of proliferation and/or differentiation of receptor-bearing cells. Based upon the tissue distribution of HEK receptors shown in Table 4, treatment with the pharmaceutical compositions of the invention may be 20 particularly indicated for disorders involving brain, heart, muscle, lung, or pancreas. However, some HEK receptors are displayed on a wide variety of tissues, so it is understood that the effects of modulating receptor activation may not be limited to those tissues described 25 herein.

The following examples are offered to more fully illustrate the invention, but are not to be construed as limiting the scope thereof. Recombinant DNA methods used in the following examples are generally as described in Sambrook et al. Molecular Cloning: A Laboratory Manual Cold Spring Harbor Laboratory Press, 2nd ed. (1989)

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EXAMPLE 1

Cloning and Sequencing of HEK Receptor cDNA

We have isolated clones for five members of 5 the EPH sub-family of receptor PTKs from a human fetal brain cDNA library. Oligonucleotides were designed based on conserved amino acid sequences within the kinase domain. Primer I was based on the amino acid sequence Trp-Thr-Ala-Pro-Glu-Ala-Ile (SEQ ID NO: 1), 10 which is well-conserved among PTKs of many families. Primer II was based on the sequence Val-Cys-Lys-Val-Ser-Asp-Phe-Gly (SEQ ID NO: 2), which is invariant among EPH sub-family members but, except for the sequence Asp-Phe-Gly, is rarely found in other PTKs. Fully degenerate oligonucleotides corresponding to reverse translations 15 of these protein sequences were synthesized and utilized as primers in a polymerase chain reaction (PCR) with disrupted phage from a human fetal brain cDNA library as the template. The products of this PCR reaction were 20 cloned into the plasmid vector pUC19 and the nucleotide sequence of the inserts was determined. Of the 35 PCR inserts sequenced, 27 were recognizable as portions of PTK genes. Their correspondence to previously published sequences is summarized in Table 1.

An asterisk indicates that different nucleic acid sequences encoded the amino acid sequence shown.

TABLE 1

Number of Clones 2	* ທ	ω	4		* 0	4
3)	4	5)	(9	(,	8	6
(SEQ ID NO: 3)	(SEQ ID NO:	(SEQ ID NO:	(SEQ ID NO:	(SEQ ID NO: 7)	(SEQ ID NO:	(SEQ ID NO: 9)
PCR Products SLGGKIPVRWTAPEAI	RGGKIPIRWTAPEAI	ALGGKIPIRWTAPEAI	RGGKIPIRWTAPEAI	GGKIPVRWTAPEAI	GAKFPIKWTAPEAI	GSTFLPLKWTAPEAI
PCR_Products VCKVSDFGLSRYLQDDTSDPTYTSSLGGKIPVRWTAPEAI	VCKVSDFGLSRVLEDDPEAAYTT	VCKVSDFGLSRFLEDDTSDPTYTSALGGKIPIRWTAPEAI	VCKVSDFGMSRVLEDDPEAAYTT	VCKVSDFGLSRVIEDDPEAVYTTT	VCKVSDFGLAR LIEDNEYTARQ	VCKVSDFGLARDIMRDSNYISK
Receptor Elk	нек4, нек7	нек5	некв	НЕК11	SRC	PDGF-β
•				-		-

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Six PCR inserts predict amino acid sequences which are identical to a portion of SRC, although they comprise two distinct nucleotide sequences. One insert appears to code for the human platelet derived growth factor (PDGF)- β receptor. The remaining 18 PCR inserts consist of 6 distinct nucleotide sequences, all of which appear to be fragments of EPH sub-family members. of the sequence predicts an amino acid sequence identical to the corresponding region of rat Elk (Lhotak 10 et al., 1991)) and is likely to represent its human homolog. Two inserts predict amino acid sequences which match the translation of the PCR fragment tyro-4 (Lai and Lemke, 1991)) but are clearly distinct at the nucleotide level while two others correspond to tyro-1 15 and tyro-5. The sixth PCR insert has a previously unreported EPH-related sequence. Since five of the clones contained portions of potential EPH sub-family members for which full-length sequences had not been reported, each was radiolabeled and used as a probe to screen a human fetal brain cDNA library. Several clones 20 corresponding to each of the five probes were isolated. For each of the five receptors, the nucleotide sequence of the clone containing the largest portion of the predicted coding region was determined.

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A single cDNA clone containing the complete coding region was isolated only for HEK4. The portions of HEK5, HEK7, HEK10 and HEK11 coding for the amino terminus of these receptors were not found in any of the clones. In order to obtain the complete coding sequence, the Rapid Amplification of cDNA Ends (RACE) technique was employed. In some cases, more than one round of RACE was necessary to obtain the missing portion of the coding region. Using this strategy, complete coding sequences were obtained for all clones except HEK7 which lacked the complete leader sequence.

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The DNA sequences of HEK5, HEK7, HEK8 and HEK11 are shown in Figures 1-4, respectively, and in SEQ ID NO: 10 (HEK5), SEQ ID NO: 12 (HEK7), SEQ ID NO: 14 (HEK 8) and SEQ ID NO: 16 (HEK11). The amino acid sequences are shown in SEQ ID NO: 11 (HEK5), SEQ ID NO: 13 (HEK7), SEQ ID NO: 15 (HEK8) and SEQ ID NO: 17 (HEK 11).

EXAMPLE 2

10 Analysis of HEK Receptor Sequences

HEK5, HEK7, HEK8 and HEK11 represent novel human EPH sub-family members, although homologs for all except HEK11 have been isolated from other species. We refer to human EPH receptor sub-family members as HEKs (human EPH-like kinases) following the nomenclature of Wicks et al., 1992). We have chosen names and numbers for these receptors to correspond with previously discovered members of the family in chicken (Ceks) and in mouse (Mek) (Sajjadi et al. 1991; Sajjadi and Pasquale, 1993; Pasquale, 1991). Extending the convention of designating the species of origin by the first letter, we refer to the rat homologs of the HEK receptors as Reks (rat EPH-like kinases).

HEK4 is the human homolog of the chicken receptor Cek4 (91% amino acid identity in the catalytic domain) and the mouse receptor Mek4 (96% amino acid identity in the catalytic domain). The amino acid sequence of HEK5 is very closely related (96% amino acid identity in the catalytic domain) to the chicken receptor Cek5 (Pasquale et al. J. Neuroscience 12, 3956-3967 (1992); Pasquale, 1991). HEK7 is probably the human homolog of the recently reported Cek7 (Sajjadi and Pasquale, 1993). HEK8 is likewise very closely related to Sek (Gilardi-Hebenstreit et al., 1992)) and Cek8 (95% amino acid identity in the catalytic domain) (Sajjadi

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and Pasquale, 1993)). The human homologs for Cek6 and Cek9 have yet to be reported, while the human homolog of Cek10 has just recently been published. One of our human receptors has no close relatives in other species and apparently represents a novel member of the EPH subfamily. We have designated this receptor HEK11, assuming that human homologs for Cek 9 and 10 will be named HEK9 and HEK10, respectively. A summary of known EPH sub-family members is shown in Table 2.

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TABLE 2 EPH receptor sub-family members

15	Human	Non-human homologs		
	ЕРН	None identified		
τ.	ECK	None identified		
	None identified#	Eek		
	HEK4*	Cek4, Mek4		
20	HEK5	Cek5, Nuk, ERK		
	None identified#	Cek6, Elk		
	HEK7	Cek7, Ehk1		
	HEK8	Cek8, Sek		
	None identified#	Cek9		
25	HEK2	Cek10		
	HEK11	None identified		
	None identified	Ehk2		

*published by Wicks et.al., 1992 as HEK

#Using the present nomenclature, the predicted human homolog of Eek is designated HEK3. For Cek6, the predicted human homolog is designated HEK6; For Cek9, the predicted human homolog is designated HEK9.

The predicted amino acid sequences of the four novel receptor clones and the previously known EPH sub-family members ECK (SEQ ID NO: 18), EPH (SEQ ID NO: 19), HEK2 (SEQ ID NO: 20) and HEK4 (SEQ ID NO: 21) were aligned as shown in Fig. 5. The four clones are closely related to each other and to the known EPH sub-family members. The extracellular domain sequences of all four novel receptors contain the Ig-loop, fibronectin-type III repeats, and cysteine-rich region characteristic of 10 EPH sub-family members. The positions of the 20 cysteine residues are conserved among all sub-family members. Also completely conserved is the portion of the catalytic domain used as the basis for the EPH sub-15 family specific primer (Val-Cys-Lys-Val-Ser-Asp-Phe-Gly, SEQ ID NO: 2, amino acids 757-764 in Fig. 5). Table 3 summarizes the percentage of sequence identity between pairs of human EPH sub-family members. The lower portion of the table shows percent amino acid identity in the catalytic domain while the upper half shows 20 percent amino acid identity in the extracellular region. The amino acid sequences of the EPH-like receptors are extremely well-conserved (60-89% amino acid identity) in the catalytic region but not as highly conserved in the 25 extracellular region (38-65% amino acid identity), as would be expected for members of the same receptor subfamily.

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TABLE 3

Eph family amino acid sequence comparison

	extracellular domains							
	EPH	ECK	HEK4	HEK5	HEK7	HEK8	HEK2	HEK11
EPH	*	47	42	38	40	43	40	42
ECK	62	*	47	41	45	46	41	46
HEK4	62	76	*	53	65	61	51	59
HEK5	60	74	81	*	52	53	63	51
HEK7	61	76	89	83	*	62	48	61
HEK8	62	76	86	85	88	*	52	57
HEK2	61	74	81	89	82	83	*	48
HEK11	60	74	83	83	85	85	80	*

Catalytic domains

Numbers shown are precent identity

10 Pairwise comparisons of amino acid sequences can be used to construct phylogenetic trees depicting the evolutionary relatedness of a family of molecules. Figure 6 is such a tree, which summarizes the relationships among the EPH sub-family members. 15 one family member is shown from each group of crossspecies homologs and the human representative was used whenever possible (refer to Table 2 for a summary of cross-species homologs). The branch lengths represent the degree of divergence between members. It has been shown previously that the EPH sub-family lies on a 20 branch evolutionarily closer to the cytoplasmic PTKs than to other receptor PTKs (Lindberg and Hunter, 1993). Interestingly, the further one moves up the tree, the more closely related the receptors become and expression becomes more localized to the brain. 25

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EXAMPLE 3

Construction and Expression of HEK Receptor Extracellular Domains

Soluble extracellular forms of HEK receptor proteins were constructed by deletion of DNA sequences encoding transmembrane and cytoplasmic domains of the receptors and introduction of a translation stop codon at the 3' end of the extracellular domain. A construct of the HEK5 extracellular domain had a stop codon introduced after lysine at position 524 as shown in Figure 1; the HEK7 extracellular domain was constructed with a stop codon after glutamine at position 547 as shown in Figure 2; the HEK 8 extracellular domain was constructed with a stop codon after threonine at position 547 as shown in Figure 3.

HEK extracellular domain was amplified from a human fetal brain cDNA library by PCR using primers 5' and 3' to the extracellular domain coding region.

For HEK5, the primers

- 5' CTGCTCGCCGCGTGGAAGAAACG (SEQ ID NO: 22) and;
- 5' GCGTCTAGATTATCACTTCTCCTGGATGCTTGTCTGGTA (SEQ ID NO: 23)

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were used to amplify the extracellular domain and to provide a restriction site for cloning into plasmid pDSR α . In addition, the following primers were used to provide a translational start site, the elk receptor signal peptide for expression; and a restriction site for cloning into pDSR α :

- 23 -

5! GCGGTCGACGCCGCCATGGCCCTGGATTGCCTGCTGTTCCTCCTG (SEQ ID NO: 24) and;

5' CGTTTCTTCCACGGCGGCGAGCAGAGATGCCAGGAGGAACAGCAGCAGCA ATC (SEQ ID NO: 25)

The resulting construct resulted in fusion of DNA encoding the elk signal sequence Met-Ala-Leu-Asp-Cys-Leu-Leu-Phe-Leu-Leu-Ala-Ser (SEQ ID NO: 26) to the first codon of the HEK5 receptor.

The resulting HEK5 extracellular domain was cloned into pDSR α after digestion with SalI and XbaI and transfected into CHO cells for expression.

HEK8 extracellular domain was amplified from a human fetal brain cDNA library by PCR using primers 5' and 3' to the extracellular domain coding region. For HEK8, the primers

- 5' GAATTCGTCGACCCGGCGAACCATGGCTGGGAT and
- 20 5' GAATTCTCTAGATTATCATGTGGAGTTAGCCCCATCTC

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were used to amplify the extracellular domain and to provide restriction sites for cloning into plasmid $pDSR\alpha$.

25 The resulting HEK8 extracellular domain was cloned into pDSR α after digestion with SalI and XbaI and transferred CHO cells for expression.

HEK7 extracellular domain was amplified from a human fetal brain cDNA library by PCR using primers 5' and 3' to the extracellular domain coding region. For HEK7, the primers

- 5'TTCGCCCTATTTTCGTGTCTCTTCGGGATTTGCGACGCTCTCCGGACCCTCCTG
- 35 5' GAATTCTCTAGATTATCACTGGCTTTGATCGCTGGAT

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were used to amplify the extracellular domain. In addition, the following primers were used to provide a translational start site, the HEK8 receptor signal peptide sequence, and restriction site for cloning into plasmid pDSR α .

- 5 '
 GAATTCGTCGACCCGGCGAACCATGGCTGGGATTTTCTATTTCGCCCTATTTTCGT
 GTCT
- 10 5' GAATTCTCTAGATTATCACTGGCTTTGATCGCTGGAT

The resulting construct resulted in fusion of DNA incoding HEK8 signal sequence Met-Ala-Gly-Ile-Phe-Tyr-Phe-Ala-Leu-Phe-Ser-Cys-Leu-Phe-Gly-Ile-Cys-Asp to the first codon of the HEK7 receptor.

The resulting HEK7 extracellular domain was cloned into pDSR α after digestion with SalI and XbaI and transfected into CHO cells for expression.

20 EXAMPLE 4

Antibodies to HEK Receptors

Antibodies to HEK receptor proteins were generated which recognize the extracellular domain by using bacterial fusion proteins as the antigen.

Antibodies were also generated which recognize the cytoplasmic domain by using synthetic peptides as the antigen.

The methodology employed has been previously described (Harlow and Lane, In <u>Antibodies: A Laboratory Manual, 1988)</u>. For the extracellular domain antibodies, cDNAs were inserted into the pATH vector (see Table 4 for the regions of each receptor encoded by this construct). These constructs were expressed in bacteria and the resultant TrpE-fusion proteins were purified by SDS-polyacrylamide gel electrophoresis. For the

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cytoplasmic domain anti-peptide antibodies, peptides were synthesized (see Table 4 for the sequences) and covalently coupled to keyhole limpet hemocyanin. The fusion proteins and coupled peptides were used as antigens in rabbits and antisera were generated and characterized as described (Harlow and Lane, 1988). Anti-peptide antibodies were affinity purified by using a SulfoLink kit (Pierce, Rockford IL).

10

TABLE 4

HEK Receptor Antigens

15 Receptor		Peptide Sequences	Amino Acids in Fusion Protein
	HEK4	CLETQSKNGPVPV	22-159
	HEK5	CRAQMNQIQSVEV	31-168
	HEK7	CMKVQLVNGMVPL	335-545
20	HEK8	CMRTQMQQMHGRMVPV	27-188
	HEK11	CQMLHLHGTGIQV	187-503

EXAMPLE 5

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HEK/TrkB Chimeric Receptors

1. Generation of pSJA1 encoding rat trkB cytoplasmic domain.

All of the chimeric receptors are composed of the extracellular domain and the transmembrane region of one of the HEK receptors and the intracellular portion of rat trkB. To simplify each individual construction, an intermediate or parental plasmid, called RtrkB/AflII (or pSJA1), was generated. First, without altering the coded peptide sequence, an AflII site (CTTAAG) was introduced into position 2021 (cytosine at position 2021

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(C2021) to guanine at position 2026 (G2026, CTCAAG) of the rat trkB cDNA (Middlemas, et al., Mol. Cell. Biol. 11, 143-153 (1991)) by PCR aided mutagenesis. Briefly, PCR primers were synthesized based on the rat trkB cDNA sequence. Primer I encompassed C2003 to G2034 of the This primer contained two mutations, a cytosine to thymine(T) substitution at position 2023 (C2023T) and an insertion of an adenine (A) in between T2013 and These mutations created the AfIII site at position C2021 and an additional XhoI site flanking the 10 AflII site. Primer II was in the reverse direction encompassing T2141 to A2165 of the cDNA which bore an Apal site. The PCR fragment produced with these primers and the rat trkB cDNA template was digested with XhoI and ApaI enzymes and sub cloned into the XhoI and ApaI 15 sites of an expression vector, pcDNA3 (InVitroGen), to generate pSJA1-b. Following, pSJA1-b was linearized with ApaI and ligated with a BanII digested rat trkB cDNA fragment (G2151 to G4697) to reconstitute a larger fragment (C2021 to G4697) including the coding sequence 20 of the whole intracellular domain of the rat trkB protein (L442 to G790) and 1571 residues (A3131 to G4697) of the 1627 nucleotide 3'-end non-coding region of the cDNA.

Generation of HEK8/rat trkB (pSJA5)

HEK8/rat trkB chimera was generated with a similar strategy as mentioned above. A SalI/BsaI cDNA fragment was first isolated from plasmid TK10/FL13. This fragment included the nucleotide sequence from the beginning to T1689 of the HEK8 cDNA (Figure 3). Then, a pair of oligonucleotides was synthesized based on the HEK8 cDNA sequence. The sequence of the first oligonucleotide was the same as G1690 to C1740 of the Hek8 cDNA, with an additional C residue added to its 3'-end. The second oligonucleotide was in the reverse

10

chimerical construct.

orientation of the HEK8 cDNA. It contained C1694 to C1740 of the HEK8 cDNA sequence and an additional five residue motif, TTAAG, at its 5'-end. These two oligonucleotides were kinased and annealed with equal molar ratio, to create a double strand DNA fragment with the sequence of G1690 to C1740 of the HEK8 cDNA and with the BsaI and the AflII cohesive ends at its 5' and 3' ends, respectively. This fragment was ligated together with the SalI/BsaI cDNA fragment into XhoI/AflII linearized pSJA1 to generate the HEK8/RtrkB (pSJA5)

3. Generation of HEK11/rat trkB (pSJA6) chimera.

To generate the HEK11/rat trkB chimera, a 15 SalI/AccI fragment covering the sequence of nucleotide C1 to T1674 of the HEK11 cDNA (Figure 4) was first isolated from plasmid TK19T3. Then, a pair of oligonucleotides was synthesized based on the HEK11 cDNA sequence. The first oligonucleotide had the same 20 sequence as from nucleotide A1666 to T1691 of the HEK11 cDNA, which contained the AccI site. The second oligonucleotide was in the reverse orientation of the HEK11 cDNA. It encompassed G1895 to T1919 of the HEK11 cDNA sequence. An additional ten residue motif, CCCGCTTAAG, was added to the 5'-end of this 25 oligonucleotide to introduce an AflII site, which would be used to link the external domain and the transmembrane region of the HEK11 receptor to the intracellular domain of the rat trkB cDNA cloned in pSJA1 in the same reading frame. PCR was performed with 30 these oligonucleotides as primers and the HEK11 cDNA as template. The PCR fragment was digested with AccI and AfIII enzymes and ligated with the SalI/AccI cDNA fragment and the XhoI/AflII linearized pSJA1 to generate the HEK11/rat trkB (pSJA6) chimerical construct. 35

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EXAMPLE 6

Tissue Distribution of HEK Receptors

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The distribution of mRNA expression for HEK4, HEK5, HEK7, HEK8 and HEK11 receptors in human and rat tissues was examined by Northern blot hybridization.

Rat total RNA was prepared from tissues using the method of Chomczynski and Sacchi (Anal. Biochem 162, 156-159 (1987)). The RNA was separated by formaldehyde-10 agarose electrophoresis and transferred to Hybond-N membranes (Amersham, Arlington Heights, IL) using 20X SSC (Maniatis et al. 1982). The membrane was dried at 80°C in vacuo for 30 minutes, then crosslinked for 3 minutes on a UV transilluminator (Fotodyne, New Berlin, 15 The membrane was prehybridized for 2 hours at 42°C in 50% formamide, 5X SSPE, 5X Denhardt's, 0.2% SDS, and 100 μg/ml denatured herring sperm DNA (Maniatis et al. 1982). Northern blots of human tissue were purchased 20 from Clontech (Palo Alto, CA). Probes were prepared by labeling the fragment of cDNA which encoded the extracellular domain of the receptor with 32p-dCTP using a hexanucleotide random priming kit (Boehringer Mannheim, Indianapolis, IN) to a specific activity of at least $1x10^9$ cpm/ug. The probe was hybridized to the 25 membrane at a concentration of 1-5 ng/ml at 42°C for 24 to 36 hours in a buffer similar to the prehybridization buffer except that 1X Denhardt's was used. After hybridization, the membranes were washed 2 times for 5 minutes each in 2X SSC, 0.1% SDS at room temperature 30 followed by two 15 minute washes in 0.5% SSC, 0.1% SDS at 55°C. Blots were exposed for 1-2 weeks using Kodak XAR film (Kodak, Rochester, NY) with a Dupont Lightning Plus intensifying screen. The results are shown in 35 Figures 7-11.

Homologs for HEK4 have been previously identified from mouse, chicken, and rat. In the adult mouse, expression is detected primarily in the brain and testis (Sajjadi et al. 1991). A slightly different pattern was found in adult chicken tissues, with the main sources of expression being the brain, liver, and Lower levels of expression were detectable in the lung and heart (Marcelle & Eichmann, Oncogene 1, 2479-2487 (1992)). A fragment of the Rek4 gene (tyro-4) has been isolated and used to look at tissue expression 10 in the adult rat (Sajjadi et al. 1991). The brain was the only tissue that expressed Rek4 mRNA. However, RNA from lung or testis were not examined. Previous studies on HEK4 only looked at the expression of the mRNA in cell lines, where it was found in one pre-B cell line 15 and two T-cell lines (Wicks et al. 1992). significance of this with regard to in vivo expression remains to be determined. In this study we have looked at the HEK4 expression in human tissues, and also the 20 expression of Rek4 in rat tissues. The HEK4 mRNA corresponds to a single transcript with a size of about 7 kb (Fig 7A). HEK4 mRNA was most abundantly expressed in placenta, with lower levels present in heart, brain, lung, and liver. On prolonged exposures, trace amounts 25 of mRNA were detectable in kidney and pancreas. Expression in the rat was more similar to that detected in the mouse and chicken. Rek4 was expressed at the lowest levels of any of the family members characterized herein. A transcript of about 7 kb was detectable in rat lung, with a lower amount detectable in brain (Fig. 30 7B). Also, a 4 kb transcript was expressed in rat testis. Because the transcripts were barely detectable using total RNA, some of the other rat tissues may contain amounts of Rek4 below the level of detection.

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Torrespond to the blots

Torrespond to 
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brain and lower expression in the heart, lung and kidney. A fragment of Rek8 (tyro-1) was used to look at expression in rat tissues, with expression found only in the brain (Lai & Lemke, 1991). We found that HEK8 mRNA was expressed at levels comparable to that of HEK5. Multiple transcripts were also observed, the most abundant at 7 kb and 5 kb. The highest level of mRNA expression was seen in the brain, although substantial levels were detected in other tissues including heart, 10 lung, muscle, kidney, placenta, and pancreas. Expression in liver was much lower than in the other The only difference in expression patterns between human and mouse was expression in human muscle, also seen for Cek8 in chicken. Among the rat tissues, 15 Rek8 was most highly expressed in the brain, followed by the lung, heart, and testis (Fig. 10B). In contrast to HEK8, expression of Rek8 appeared to be lower in muscle and kidney, two tissues where HEK8 was readily detectable. In addition, Rek8 was not expressed as a 20 5.0 kb transcript, as it was not visible even on prolonged exposures.

During the analysis of this family, we deduced that HEK7 is the human homolog of Cek7. The only 25 expression seen in adult chicken was an 8.5 kb transcript found in the brain (Sajjadi & Pasquale, 1993). Of the five EPH sub-family members described here, HEK7 was the most restricted in its expression pattern. Analysis of human mRNA revealed significant 30 expression only in the brain, with a much lower level detectable in the placenta (Fig. 9A). Prolonged exposures did not reveal expression in any other tissue examined. Two prominent transcripts were found in brain, the most highly expressed with a size of 6 kb and the other with a length of 9 kb. In the placenta, 35 however, only the 9 kb transcript was detected. Rek7

mRNA was expressed in a pattern similar to HEK7. The highest level of expression was found in brain, with a much lower level in ovary (Fig. 9B). The transcripts were of similar size as for HEK7, with the 6 kb transcript detected only in brain.

HEK11 was expressed as several transcripts, with major mRNAs of length 7.5, 6.0 and 3.0 kb and minor transcripts of 4.4 and 2.4 kb (Fig. 11A). All five

10 mRNAs were expressed at the highest levels in brain, followed by heart. Placenta, lung and kidney had significant amounts of four of the five transcripts, with lower expression seen in muscle. Pancreas had barely detectable amounts of HEK11 mRNA, while liver had no detectable HEK11 transcript. Rek11 had a similar pattern of expression, with four transcripts (10, 7.5, 3.5 and 3.0 kb) detected in brain (Fig. 11B).

The relative level of mRNA expression for each of the five receptors in all tissues studied is summarized in Table 5.

HEK5

HEK4

 $^{\rm -}$ 33 $^{\rm -}$ Table 5 Tissue Distribution of HEK Receptors

HEK7

HEK8

,	Brain	++	++	++	+++	++
	Heart	+	+	bd	++	+
	Kidney	+	+ ,	bd	+	+
	Liver	+	+	bd .	+	bd
	Lung	+	+	bd	++	+
	Muscle	+	+	bd	++	+
	Pancreas	+	++	bd	+	bd
	Placenta	+++	+++	bd	++	+
5						
	Rat	HEK4	HEK5	HEK7	HEK8	HEK11
	Brain	+	++	+++	+++	++
	Heart	bd	bd	bd	+	bd
	Intestine	bd	+++	bd	bd	bd
	Kidney	bd	++	bd	bd	bd
	Liver	bd	bd	bd	bd	bd .
	Lung	+	+	bd	++	bd
	Muscle	bd	bd	bd	bd	bd
	Ovary	bd	+	+	· bd /	bd
	Stomach	bd	+	bd	bd	bd
	Testis	+	bd	bd	+	bd
	Thymus	bd .	+	bd	bd	bd

bd= below detection

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The transcripts for HEKs 4,5,8, and 11 were rather widely distributed in human tissue while HEK7 was specific for brain. Expression patterns between rat and human tissue were roughly comparable given that the rat blots were less sensitive due to the use of total RNA rather than polyA⁺. As was found for the Cek mRNAs by Sajjadi and Pasquale (Sajjadi & Pasquale, 1993), often there were several different size transcripts detected for a single receptor. The size distribution of the transcripts appears to be both tissue and species specific. Previous work has shown that the smaller transcript of Mek4 encodes a potentially secreted receptor (Sajjadi et al. 1991).

The following sections describe Materials and Methods used to carry out experiments described in Example 1.

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Isolation, cloning and sequencing of HEK receptor cDNAs

Fragments containing a portion of the catalytic domain of EPH sub-family receptors were generated using a polymerase chain reaction (PCR) with disrupted phage from a human fetal brain cDNA library as a template. A 10µl aliquot of the cDNA library (Stratagene, La Jolla, CA) was treated at 70°C for 5 minutes to disrupt the phage particles, then cooled on wet ice. The disrupted phage were added to 10µl of 10x Tag polymerase buffer, 8ul of 2mM each dNTP, 100 picomoles of each primer, and 1.5 μ l of Tag polymerase (Promega, Madison, WI) in a total volume of 100μl. reaction was run for 35 cycles, each consisting of 1 minute at 96°C, 1 minute at 50°C, and 2 minutes at 72°C. A 5 minute, 72°C incubation was added at the end to ensure complete extension. The primers used were degenerate mixtures of oligonucleotides based on amino

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acid sequences which are highly conserved among EPH sub-family members.

5'AGGGAATTCCAYCGNGAYYTNGCNGC' (SEQ ID NO: 27); 5'AGGGGATCCRWARSWCCANACRTC'(SEQ ID NO: 28).

The products of the PCR reaction were digested with EcoRI and BamHI and cloned into M13mp19 (Messing, Methods Enzymol. (1983)) for sequence analysis. five clones which were identified as fragments of EPH 10 receptor sub-family members were labeled with 32p-dCTP by random priming and each was used to screen Genescreen nitrocellulose filters (NEN, Boston, MA) containing plaques from the human fetal brain cDNA library. Phage 15 stocks prepared from positively screening plagues were plated and rescreened with the same probe in order to obtain single clones. cDNA inserts were transferred into pBluescript using the in vivo excision protocol supplied with the cDNA library (Stratagene, La Jolla, 20 CA). Nucleotide sequences were determined using Tag DyeDeoxy Terminator Cycle Sequencing kits and an Applied Biosystems 373A automated DNA sequencer (Applied Biosystems, Foster City, CA).

25 <u>5' Race</u>

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The 5' ends of the cDNAs were isolated using a 5' RACE kit (GIBCO/BRL, Gaithersburg, MD) following the manufacturer's instructions. Excess primers were removed after first strand cDNA synthesis using ultrafree-MC cellulose filters (30,000 molecular weight cutoff, Millipore, Bedford, MA). Amplified PCR products were digested with the appropriate restriction enzymes, separated by agarose gel electrophoresis, and purified using a Geneclean kit (Bio101, La Jolla, CA). The purified PCR product was ligated into the plasmid vector pUC19 (Yanisch-Perron et al. Gene 33, 103-119 (1985))

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which had been digested with appropriate restriction enzymes and the ligation mixture was introduced into host bacteria by electroporation. Plasmid DNA was prepared from the resulting colonies. Those clones with the largest inserts were selected for DNA sequencing.

While the present invention has been described in terms of preferred embodiments, it is understood that variations and modifications will occur to those skilled in the art. Therefore, it is intended that the appended claims cover all such equivalent variations which come within the scope of the invention as claimed.

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SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: Amgen Inc.
 - (ii) TITLE OF INVENTION: EPH-Like Receptor Protein Tyrosine Kinases
 - (iii) NUMBER OF SEQUENCES: 28
 - (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Amgen Patent Operations/RBW
 - (B) STREET: 1840 Dehavilland Drive
 - (C) CITY: Thousand Oaks
 - (D) STATE: California
 - (E) COUNTRY: USA
 - (F) ZIP: 91320
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Winter, Robert B.
 - (C) REFERENCE/DOCKET NUMBER: A-287
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 7 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

Trp Thr Ala Pro Glu Ala Ile

- (2) INFORMATION FOR SEQ ID NO:2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Val Cys Lys Val Ser Asp Phe Gly
1 5

- (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 40 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: \single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Val Cys Lys Val Ser Asp Phe Gly Leu Ser Arg Tyr Leu Gln Asp Asp 1 5 10 15

Thr Ser Asp Pro Thr Tyr Thr Ser Ser Leu Gly Gly Lys Ile Pro Val 20 25 30

Arg Trp Thr Ala Pro Glu Ala Ile 35 40

- (2) INFORMATION FOR SEQ ID NO:4:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 38 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Val Cys Lys Val Ser Asp Phe Gly Leu Ser Arg Val Leu Glu Asp Asp 1 5 10 15

Pro Glu Ala Ala Tyr Thr Thr Arg Gly Gly Lys Ile Pro Ile Arg Trp

Thr Ala Pro Glu Ala Ile 35

- (2) INFORMATION FOR SEQ ID NO:5:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 40 amino acids

 - (B) TYPE: amino acid(C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Val Cys Lys Val Ser Asp Phe Gly Leu Ser Arg Phe Leu Glu Asp Asp

Thr Ser Asp Pro Thr Tyr Thr Ser Ala Leu Gly Gly Lys Ile Pro Ile

Arg Trp Thr Ala Pro Glu Ala Ile 35

- (2) INFORMATION FOR SEQ ID NO:6:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 38 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Val Cys Lys Val Ser Asp Phe Gly Met Ser Arg Val Leu Glu Asp Asp

Pro Glu Ala Ala Tyr Thr Thr Arg Gly Gly Lys Ile Pro Ile Arg Trp 20

Thr Ala Pro Glu Ala Ile 35

- (2) INFORMATION FOR SEQ ID NO:7:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 38 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: -

Val Cys Lys Val Ser Asp Phe Gly Leu Ser Arg Val Ile Glu Asp Asp 1 5. 10 15

Pro Glu Ala Val Tyr Thr Thr Thr Gly Gly Lys Ile Pro Val Arg Trp 20 25 30

Thr Ala Pro Glu Ala Ile 35

- (2) INFORMATION FOR SEQ ID NO:8:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 36 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Val Cys Lys Val Ser Asp Phe Gly Leu Ala Arg Leu Ile Glu Asp Asn

Glu Tyr Thr Ala Arg Gln Gly Ala Lys Phe Pro Ile Lys Trp Thr Ala
20 25 30

Pro Glu Ala Ile 35

- (2) INFORMATION FOR SEQ ID NO:9:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 37 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein

										_							
								EQ I									
	Val 1	. Суз	Lys	Val	Ser 5	: Asp	Phe	: Gly	Leu	Ala 10	Arg	J Asp	Ile	Met	Arg 15	Asp	
	Ser	: Азп	туг	11e 20	e Ser	Lys	Gly	Ser	Thr 25	Phe	e Leu	Pro	Leu	Lys 30	Trp	Thr	
	Ala	Pro	Glu 35	ı Ala	ılle	;	-		**								
(2)	INFO	RMAI	ON	FOR	SEQ	ID N	10:10	:									
	(i)	(E (C	A) LE B) TY C) SI	NGTH	nucl	62 t eic SS:	acid	pair l	:s								X.
	(ii)	MOI	ECUI	E TY	PE:	CDNA	7										
	(ix)		1) NA	: AME/F CAT			2913										
	(xi)	SEÇ	QUENC	CE DE	ESCRI	PTIC	N: S	SEQ I	D NC	:10:							
													ACA Thr				48
													GAA Glu 30				96
													CAG Gln				144
													AAG Lys				192
													TTT Phe				240
													AAG Lys				288
TTC Phe	AAC. Asn	CTC Leu	TAT Tyr 100	TAC Tyr	TAT Tyr	GAG Glu	GCT Ala	GAC Asp 105	TTT Phe	GAC Asp	TCG Ser	GCC Ala	ACC Thr 110	AAG Lys	ACC Thr		336

						GAT Asp 125		GCA Ala	384
						CGC Arg			432
						CGC Arg		_	480
						CTC Leu			528
						AAT Asn			576
						CTG Leu 205			624
						GTA Val			672
						ATC Ile			720
						ACC Thr			768
						GAT Asp			816
						GGG Gly 285			864
						GAC Asp			912
						GTG Val			960
						CCC Pro			1008

												AAG Lys				1	056
												GTA Val 365				1	.104
												ATC Ile				1	.152
												GTG Val				1	.200
												GTG Val				1	248
Thr	Asn	Gln	Ala 420	Ala	Pro	Ser	Ala	Val 425	Ser	Ile	Met	CAT His	Gln 430	Val	Ser	1	296
Arg	Thr	Val 435	Asp	Ser	Ile	Thr	Leu 440	Ser	Trp	Ser	Gln	CCG Pro 445	Asp	Gln	Pro	. 1	344
Asn	Gly 450	Val	Ile	Leu	Asp	Tyr 455	Glu	Leu	Gln	Tyr	Tyr 460	GAG Glu	Lys	Glu	Leu	1	392
Ser 465	Glu	Tyr	Asn	Ala	Thr 470	Ala	Ile	Lys	Ser	Pro 475	Thr	AAC Asn	Thr	Val	Thr 480	1	440
Gly	Leu	Lys	Ala	Gly 485	Ala	Ile	Tyr	Val	Phe 490	Gln	Val	CGG Arg	Ala	Arg 495	Thr	1	488
Val	Ala	Gly	Tyr 500	Gly	Arg	Tyr	Ser	Gly 505	Lys	Met	Tyr	TTC Phe	Gln 510	Thr	Met	1	536
Thr	Glu	Ala 515	Glu	Tyr	Gln	Thr	Ser 520	Ile	Gln	Glu	Lys	TTG Leu 525	Pro	Leu	Ile	1	584
Ile	Gly 530	Ser	Ser	Ala	Ala	Gly 535	Leu	Val	Phe	Leu	Ile 540	GCT Ala	Val	Val	Val	1	632
ATC Ile 545	GCC Ala	ATC Ile	GTG Val	TGT Cys	AAC Asn 550	AGA Arg	CGG Arg	GGG Gly	TTT Phe	GAG Glu 555	CGT Arg	GCT Ala	GAC Asp	TCG Ser	GAG Glu . 560	1	680

TAC Tyr	ACG Thr	GAC Asp	AAG Lys	CTG Leu 565	CAA Gln	CAC His	TAC Tyr	ACC Thr	AGT Ser 570	GGC Gly	CAC His	ATA Ile	ACC Thr	CCA Pro 575	GGC Gly	· ·	1728
	AAG Lys																1776
	CGG [.] Arg																1824
	GTG Val 610																1872
	CTG Leu																1920
	GGC Gly																1968
-	ATG Met																2016
	ACC Thr																2064
	TCC Ser 690																2112
	CAG Gln								_	_							2160
	GCA Ala																2208
	GTC Val																2256
	TTT Phe																2304
	GGA Gly 770																2352

CGG Arg 785	AAG Lys	TTC Phe	ACC Thr	TCG Ser	GCC Ala 790	AGT Ser	GAT Asp	GTG Val	TGG Trp	AGC Ser 795	TAC Tyr	GGC Gly	ATT Ile	GTC Val	ATG Met 800	2400
												GAC Asp				2448
												CTG Leu				2496
												GAC Asp 845				2544
												GTC Val				2592
-												ATG Met				2640
												ATC Ile				2688
												ATC Ile				2736
												TCC Ser 925				2784
												GGG Gly				2832
												GTG Val				2880
			CAG Gln							TGA	CATTO	CAC (CTGC	CTCGO	SC	2930
TCAC	CTC	rtc (CTCC	AAGC	cc co	GCCC	CTC	r GC								2962

(2) INFORMATION FOR SEQ ID NO:11:

- (i) SEQUENCE CHARACTERISTICS:

 (A) LENGTH: 970 amino acids

 (B) TYPE: amino acid

 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Leu Leu Ala Ala Val Glu Glu Thr Leu Met Asp Ser Thr Thr Ala Thr Ala Glu Leu Gly Trp Met Val His Pro Pro Ser Gly Trp Glu Glu Val Ser Gly Tyr Asp Glu Asn Met Asn Thr Ile Arg Thr Tyr Gln Val Cys Asn Val Phe Glu Ser Ser Gln Asn Asn Trp Leu Arg Thr Lys Phe Ile Arg Arg Arg Gly Ala His Arg Ile His Val Glu Met Lys Phe Ser Val 70 Arg Asp Cys Ser Ser Ile Pro Ser Val Pro Gly Ser Cys Lys Glu Thr Phe Asn Leu Tyr Tyr Glu Ala Asp Phe Asp Ser Ala Thr Lys Thr Phe Pro Asn Trp Met Glu Asn Pro Trp Val Lys Val Asp Thr Ile Ala Ala Asp Glu Ser Phe Ser Gln Val Asp Leu Gly Gly Arg Val Met Lys 135 Ile Asn Thr Glu Val Arg Ser Phe Gly Pro Val Ser Arg Ser Gly Phe Tyr Leu Ala Phe Gln Asp Tyr Gly Gly Cys Met Ser Leu Ile Ala Val Arg Val Phe Tyr Arg Lys Cys Pro Arg Ile Ile Gln Asn Gly Ala Ile Phe Gln Glu Thr Leu Ser Gly Ala Glu Ser Thr Ser Leu Val Ala Ala Arg Gly Ser Cys Ile Ala Asn Ala Glu Glu Val Asp Val Pro Ile Lys 215 Leu Tyr Cys Asn Gly Asp Gly Glu Trp Leu Val Pro Ile Gly Arg Cys Met Cys Lys Ala Gly Phe Glu Ala Val Glu Asn Gly Thr Val Cys Arg 250 Gly Cys Pro Ser Gly Thr Phe Lys Ala Asn Gln Gly Asp Glu Ala Cys

Thr His Cys Pro Ile Asn Ser Arg Thr Thr Ser Glu Gly Ala Thr Asn 275 280 285

Cys	Val 290	Суз	Arg	Asn	Gly	Tyr 295	Tyr	Arg	Ala	Asp	Leu 300	Asp	Pro	Leu	Asp
Met 305	Pro	Cys	Thr	Thr	Ile 310	Pro	Ser	Ala	Pro	Gln 315	Ala	Val	Ile	Ser	Ser 320
Val	Asn	Glu	Thr	Ser 325	Leu	Met	Leu	Glu	Trp 330	Thr	Pro	Pro	Arg	Asp 335	Ser
Gly	Gly	Arg	Glu 340	Asp	Leu	Val	Tyr	Asn 345	Ile	Ile	Суз	Lys	Ser 350	Суз	Gly
Ser	Gly	Arg 355	Gly	Ala	Суз	Thr	Arg 360	Суз	Gly	Asp	Asn	Val 365	Gln	Tyr	Ala
Pro	Arg 370	Gln	Leu	Gly	Leu	Thr 375	Glu	Pro	Arg	Ile	Tyr 380	Ile	Ser	Asp	Leu
Leu 385	Ala	His	Thr	Gln	Tyr 390	Thr	Phe	Glu	Ile	Gln 395	Ala	Val	Asn	Gly	Val 400
Thr	Asp	Gln	Ser	Pro 405	Phe	Ser	Pro	Gln	Phe 410	Ala	Ser	Val	Asn	Ile 415	Thr
Thr	Asn	Gln	Ala 420	Ala	Pro	Ser	Ala	Val 425	Ser	Ile	Met	His	Gln 430	Val	Ser
Arg	Thr	Val 435	Asp	Ser	Ile	Thr	Leu 440	Ser	Trp	Ser	Gln	Pro 445	Asp	Gln	Pro
Asn	Gly 450	Val	Ile	Leu	Asp	Туг 455	Glu	Leu	Gln	туг	Туг 460	Glu	Lys	Glu	Leu
Ser 465	Glu	Tyr	Asn	Ala	Thr 470	Ala	Ile	Lys	Ser	Pro 475	Thr	Asn	Thr	Val	Thr 480
Gly	Leu	Lys	Ala	Gly 485	Ala	Ile	Tyr	Val	Phe 490	Gln	Val	Arg	Ala	Arg 495	Thr
		_	Tyr 500	_		_		505	-		-		510		
		515	Glu				520					525			
Ile	Gly 530	Ser	Ser	Ala	Ala	Gly 535	Leu	Val	Phe	Leu	11e 540	Ala	Val	Val	Val
545			Val		550					555					560
			Lys	565					570					575	
Met	Lys	Ile	Tyr 580	Ile	Asp	Pro	Phe	Thr 585	Tyr	Glu	Asp	Pro	Asn 590	Glu	Ala

Val Arg Glu Phe Ala Lys Glu Ile Asp Ile Ser Cys Val Lys Ile Glu 600 Gln Val Ile Gly Ala Gly Glu Phe Gly Glu Val Cys Ser Gly His Leu Lys Leu Pro Gly Lys Arg Glu Ile Phe Val Ala Ile Lys Thr Leu Lys Ser Gly Tyr Thr Glu Lys Gln Arg Arg Asp Phe Leu Ser Glu Ala Ser 650 645 Ile Met Gly Gln Phe Asp His Pro Asn Val Ile His Leu Glu Gly Val Val Thr Lys Ser Thr Pro Val Met Ile Ile Thr Glu Phe Met Glu Asn Gly Ser Leu Asp Ser Phe Leu Arg Gln Asn Asp Gly Gln Phe Thr Val Ile Gln Leu Val Gly Met Leu Arg Gly Ile Ala Ala Gly Met Lys Tyr Leu Ala Asp Met Asn Tyr Val His Arg Asp Leu Ala Ala Arg Asn Ile 725 Leu Val Asn Ser Asn Leu Val Cys Lys Val Ser Asp Phe Gly Leu Ser Arg Phe Leu Glu Asp Asp Thr Ser Asp Pro Thr Tyr Thr Ser Ala Leu 760 Gly Gly Lys Phe Pro Ile Arg Trp Thr Ala Pro Glu Ala Ile Gln Tyr Arg Lys Phe Thr Ser Ala Ser Asp Val Trp Ser Tyr Gly Ile Val Met Trp Glu Val Met Ser Tyr Gly Glu Arg Pro Tyr Trp Asp Met Thr Asn 810 Gln Asp Val Ile Asn Ala Ile Glu Gln Asp Tyr Arg Leu Pro Pro Pro 825 Met Asp Cys Pro Ser Ala Leu His Gln Leu Met Leu Asp Cys Trp Gln 840 Lys Asp Arg Asn His Arg Pro Lys Phe Gly Gln Ile Val Asn Thr Leu Asp Lys Met Ile Arg Asn Pro Asn Ser Leu Lys Ala Met Ala Pro Leu Ser Ser Gly Ile Asn Leu Pro Leu Leu Asp Arg Thr Ile Pro Asp Tyr 890 885

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Thr	Ser	Phe	Asn 900	Thr	Val	Asp	Glu	Trp 905	Leu	Glu	Ala	Ile	Lys 910	Met	GJŽ
Gln	Tyr	Lys 915	Glu	Ser	Phe	Ala	Asn 920	Ala	Gly	Phe	Thr	Ser 925	Phe	Asp	Val
Val	Ser 930	Gln	Met	Met	Met :	Glu 935	Asp	Ile	Leu	Arg	Val 940	Gly	Val	Thr	Lev
Ala 945	Gly	His	Gln	Lys	Lys 950	Ile	Leu	Asn	Ser	Ile 955	Gln	Val	Met	Arg	Ala 960
Gln	Met	Asn	Gln	Ile 965	Gln	Ser	Val	Glu	Val 970						

(2) INFORMATION FOR SEQ ID NO:12:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 3162 base pairs (B) TYPE: nucleic acid

 - (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..2976
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

		GCC Ala 5							48
		CTC Leu					 		96
		GTG Val				_	 		144
		GCT Ala							192
		TAT Tyr							240
		CAG Gln 85					 		288

								GAC Asp	336
	AGC Ser 115								384
	TAC Tyr								432
	TAC Tyr								480
	GAT Asp			_					528
	GGA Gly								576
	GCT Ala 195								624
	TCT Ser								672
	GAT Asp								720
	GTG Val								768
	CTG Leu								816
	AAT Asn 275								864
	CAC His								912
	GAA Glu								960

												CCC Pro					1008
												GTC Val					1056
												GTG Val 365					1104
												TGT Cys					1152
												CTG Leu				-	1200
												TAT Tyr					1248
Ile	Glu	Ala	Val 420	Asn	Gly	Val	Ser	Asp 425	Leu	Ser	Pro	GGA Gly	Ala 430	Arg	Gln		1296
												CCA Pro 445					1344
							_	_				ATC Ile					1392
												GAG Glu					1440
Lys	His	Phe	Glu	Lys 485	Asp	Gln	Glu	Thr	Ser 490	Tyr	Thr	ATT	Ile	Lys 495	Ser		1488
												GCT Ala					1536
												GGT Gly 525					1584
												GCA Ala					1632

CAA Gln 545	AGC Ser	CAG Gln	ATT Ile	CCT Pro	GTA Val 550	ATT Ile	GCT Ala	GTG Val	TCT Ser	GTG Val 555	ACA Thr	GTA Val	GGA Gly	GTC Val	ATT Ile 560	1680
														TGT Cys 575		1728
TAC Tyr	AGC Ser	AAA Lys	GCA Ala 580	AAA Lys	CAA Gln	GAT Asp	CCA Pro	GAA Glu 585	GAG Glu	GAA Glu	AAG Lys	ATG Met	CAT His 590	TTT Phe	CAT His	1776
														CCA Pro		1824
														GAG Glu		1872
														GAA Glu		1920
														GAA Glu 655		1968
														CAA Gln		2016
														CAT His		2064
														GTG Val		2112
														TTG Leu	AAG Lys 720	2160
AAA Lys	AAC Asn	GAT Asp	GGG Gly	CAG Gln 725	TTC Phe	ACT Thr	GTG Val	ATT Ile	CAG Gln 730	CTT Leu	GTT Val	GGC Gly	ATG Met	CTG Leu 735	AGA Arg	2208
														GTG Val		2256
														GTG Val		2304

															_	
			GAC Asp													2352
			ACC Thr													2400
			ATA Ile													2448
			ATA Ile 820													2496.
			ATG Met													2544
			CCA Pro													2592
			TGC Cys													2640
			AAC Asn													2688
			GTT Val 900													2736
			CTA Leu													2784
			AAG Lys													2832
			ATG Met													2880
			GTG Val													2928
		GAA	ATG	AAG	GTG	CAG	CTG	GTA	AAC	GGA	ATG	GTG	CCA	TTG	TAACTTCA	rG
2983 Leu			Met 980	Lys	Val	Gln	Leu	Val 985	Asn	Gly	Met	Val	Pro 990	Leu		
TAAA	TGTC	GC 7	TTTT	CAAC	T GA	ATGA	TTCI	GCA	CTTT	GTA	AACA	GCAC	TG A	GATI	TATTT	3043

3103

TAAC	CAAAF	AA. A	KGGGG	GAA	AA GO	AAAO	IACAC	ı TGF	7111	JIMM	ACC.	IAGI	ww r	CAI.	116661
CAGO	CACA	AGA A	ATTTG	TAAT	ra ot	GGTT	CATT	TG#	AGT	ATCC	AGTT	CTT	AGT (CTT	AGTCT
(2)	INFO	RMAT	ON	FOR	SEQ	ID 1	10:13	3:				*			
	- ((i) S	(A) (B)	LE)	CHAP NGTH: PE: &	991 mino	l ami	ino a		5					•
	i)	li) N	OLEC	ULE	TYPE	E: pi	cotei	in							
	()	(i) S	EQUE	ENCE	DESC	CRIPT	NOI!	SEÇ	DID	NO:1	13:				
Pro 1	Ala	Ser	Leu	Ala 5	Gly	Cys	Tyr	Ser	Ala 10	Pro	Arg	Arg	Ala	Pro 15	Leu
Trp	Thr	Cys	Leu 20	Leu	Leu	Cys	Ala	Ala 25	Leu	Arg	Thr	Leu	Leu 30	Ala	Ser
Pro	Ser	Asn 35	Glu	Val	Asn	Leu	Leu 40	Asp	Ser	Arg	Thr	Val 45	Met	Gly	Asp .
Leu	Gly 50	Trp	Ile	Ala	Phe	Pro 55	Lys	Asn	Gly	Trp	Glu 60	Glu	Ile	Gly	Glu
Val 65	Asp	Glu	Asn	Туг	Ala 70	Pro	lle	His	Thr	Туг 75	Gln	Val	Cys	Lys	Val 80
Met	Glu	Gln	Asn	Gln 85	Asn	Asn	Trp	Leu	Leu 90	Thr	Ser	Trp	Ile	Ser 95	Asn
Glu	Gly	Ala	Ser 100	Arg	Ile	Phe	Ile	Glu 105	Leu	Lys	Phe	Thr	Leu 110	Arg	Asp
Cys	Asn	Ser 115	Leu	Pro	Gly	Gly	Leu 120	Gly	Thr	Cys	Lys	Glu 125	Thr	Phe	Asn
Met	Tyr 130	Tyr	Phe	Glu	Ser	Asp 135	Asp	Gln	Asn	Gly	Arg 140	Asn	Ile	Lys	Glu
Asn 145	Gln	Tyr	Ile	Lys	Ile 150	Asp	Thr	Ile	Ala	Ala 155	Asp	Glu	Ser	Phe	Thr 160
Glu	Leu	Asp	Leu	Gly 165	Asp	Arg	Val	Met	Lys 170	Leu	Asn	Thr	Glu	Val 175	Arg
Asp	Val	Gly	Pro 180	Leu	Ser	Lys	Lys	Gly 185	Phe	Tyr	Leu	Ala	Phe 190	Gln	Asp
Val	Gly	Ala 195	Cys	Ile	Ala	Leu	Val 200	Ser	Val	Arg	Val	Tyr 205	Tyr	Lys	Lys
Cys	Pro 210	Ser	Val	Val	Arg	His 215	Leu	Ala	Val	Phe	Pro 220	Asp	Thr	Ile	Thr

Gly 225	Ala	Asp	Ser	Ser	Gln 230	Leu	Leu	Glu	Val	Ser 235	Gly	Ser	Суз	Val	Asn 240
His	Ser	Val	Thr	Asp 245	Glu	Pro	Pro	ГÀЗ	Met 250	His	Суз	Ser	Ala	Glu 255	Gly
Glu	Trp	Leu	Val 260	Pro	Ile	Gly	Lys	Cys 265	Met	Cys	Lys	Ala	Gly 270	Tyr	Glu
Glu	Lys	Asn 275	Gly	Thr	Cys	Gln	Val 280	Суѕ	Arg	Pro	Gly	Phe 285	Phe	Lys	Ala
Ser	Pro 290	His	Ile	Gln	Ser	Суз 295	Gly	Lys	Cys	Pro	Pro 300	His	Ser	Tyr	Thr
His 305	Glu	Glu	Ala	Ser	Thr 310	Ser	Суѕ	Val	Cys	Glu 315	Lys	Asp	Tyr	Phe	Arg 320
Arg	Glu	Ser	Asp	Pro 325	Pro	Thr	Met	Ala	Cys 330	Thr	Arg	Pro	Pro	Ser 335	Ala
Pro	Arg	Asn	Ala 340	Ile	Ser	Asn	Val	Asn 345	Glu	Thr	Ser	V al	Phe 350	Leu	Glu
Trp	Ile	Pro 355	Pro	Ala	Asp	Thr	Gly 360	Gly	Arg	Lys	Asp	Val 365	Ser	Tyr	Tyr
Ile	Ala 370	Cys	Lys	Lys	Суѕ	Asn 375	Ser	His	Ala	Gly	Val 380	Суз	Glu	Glu	Cys
Gly 385	Gly	His	Val	Arg	Tyr 390	Leu	Pro	Arg	Gln	Ser 395	Gly	Leu	Lys	Asn	Thr 400
Ser	Val	Met	Met	Val 405	Asp	Leu	Leu	Ala	His 410	Thr	Asn	Tyr	Thr	Phe 415	Glu
Ile	Glu	Ala	Val 420	Asn	Gly	Val	Ser	Asp 425	Leu	Ser	Pro	Gly	Ala 430	Arg	Gln
Tyr	Val	Ser 435	Val	Asn	Val	Thr	Thr 440	Asn	Gln	Ala	Ala	Pro 445	Ser	Pro	Val
	Asn 450	Val	Lys	Lys	Gly	Lys 455	Ile	Ala	Lys	Asn	Ser 460	Ile	Ser	Leu	Ser
Trp 465	Gln	Glu	Pro	Asp	Arg 470	Pro	Asn	Gly	Ile	Ile 475	Leu	Glu	Tyr	Glu	Ile 480
Lys	His	Phe	Glu	Lys 485	Asp	Gln	Glu	Thr	Ser 490	Tyr	Thr	Ile	Ile	Lys 495	Ser
Lys	Glu	Thr	Thr 500	Ile	Thr	Ala	Glu	Gly 505	Leu	Lys	Pro	Ala	Ser 510	Val	Tyr
Val	Phe	Gln 515	Ile	Arg	Ala	Arg	Thr 520	Ala	Ala	Gly	Tyr	Gly 525	Val	Phe	Ser

Arg	Arg 530	Phe	Glu	Phe	Ğlu	Thr 535	Thr	Pro	Val	Phe	Ala 540	Ala	Ser	Ser	Asp
Gln 545	Ser	Gln	Ile	Pro	Val 550	Ile	Ala	Val	Ser	Val 555	Thr	Val	Gly	Val	Ile 560
Leu	Leu	Ala	Val	Val 565	Ile	Gly	Val	Leu	Leu 570	Ser	Gly	Arg	Arg	Cys 575	Gly
Tyr	Ser	Lys	Ala 580	Lys	Gln	Asp	Pro	Glu 585	Glu	Glu	Lys	Met	His 590	Phe	His
Asn	Gly	His 595	Ile	Lys	Leu	Pro	Gly 600	Val	Arg	Thr	Tyr	Ile 605	Asp	Pro	His
Thr	Tyr 610	Glu	Asp	Pro	Asn	Gln 615	Ala	Val	His	Glu	Phe 620	Ala	Lys	Glu	Ile
Glu 625	Ala	Ser	Суз	Ile	Thr 630	Ile	Glu	Arg	Val	Ile 635	Gly	Ala	Gly	Glu	Phe 640
Gly	Glu	Val	Суз	Ser 645	Gly	Arg	Leu	Lys	Leu 650	Pro	Gly	Lys	Arg	Glu 655	Leu
Pro	Val	Ala	11e 660	Lys	Thr	Leu	Lys	Val 665	Gly	Туг	Thr	Glu	Lys 670	Gln	Arg
Arg	Asp	Phe 675	Leu	Gly	Glu	Ala	Ser 680	Ile	Met	Gly	Gln	Phe 685	Asp	His	Pro
Asn	11e 690	Ile	His	Leu	Glu	Gly 695	Val	Val	Thr	Lys	Ser 700	Lys	Pro	Val	Met
Ile 705	Val	Thr	Glu	Tyr	Met 710	Glu	Asn	Gly	Ser	Leu 715	Asp	Thr	Phe	Leu	Lys 720
Lys	Asn	Asp	Gly	Gln 725	Phe	Thr	Val	Ile	Gln 730	Leu	Val	Gly	Met	Leu 735	Arg
Gly	Ile	Ser	Ala 740	Gly	Met	Lys	Tyr	Leu 745	Ser	Asp	Met	Gly	Tyr 750	Val	His
Arg	Asp	Leu 755	Ala	Ala	Arg	Asn	11e 760	Leu	Ile	Asn	Ser	Asn 765	Leu	Val	Cys
Lys	Val 770	Ser	Asp	Phe	Gly	Leu 775	Ser	Arg	Val	Leu	Glu 780	Asp	Asp	Pro	Glu
Ala 785	Ala	Tyr	Thr	Thr	Arg 790	Gly	Gly	Lys	Ile	Pro 795	Ile	Arg	Trp	Thr	Ala 800
Pro	Glu	Ala	Ile	Ala 805	Phe	Arg	Lys	Phe	Thr 810	Ser	Ala	Ser	Asp	Val 815	Trp
Ser	Tyr	Gly	Ile 820	Val	Met	Trp	Glu	Val 825	Val	Ser	Tyr	Gly	Glu 830	Arg	Pro

Tyr	Trp	Glu 835	Met	Thr	Asn	Gln	Asp 840	Val	Ile	Lys	Ala	Val 845	Glu	Glu	Gly
Tyr	Arg 850	Leu	Pro	Ser	Pro	Met 855	Asp	Суз	Pro	Ala	Ala 860	Leu	Tyr	Gln	Leu
Met 865	Leu	Asp	Cys	Trp	Gln 870	Lys	Glu	Arg	Asn	Ser 875	Arg	Pro	Lys	Phe	Asp 088
Glu	Ile	Val	Asn	Met 885	Leu	Asp	Lys	Leu	11e 890	Arg	Asn	Pro	Ser	Ser 895	Leu
Lys	Thr	Leu	Val 900	Asn	Ala	Ser	Суз	Arg 905	Val	Ser	Asn	Leu	Leu 910	Ala	Glu
His	Ser	Pro 915	Leu	Gly	Ser	Gly	Ala 920	Tyr	Arg	Ser	Val	Gly 925	Glu	Trp	Leu
Glu	Ala 930	Ile	Lys	Met	Gly	Arg 935	Tyr	Thr	Glu	Ile	Phe 940	Met	Glu	Asn	Gly
Tyr 945	Ser	Ser	Met	Asp	Ala 950	Val	Ala	Gln	Val,	Thr 955	Leu	Glu	Asp	Leu	Arg 960
Arg	Leu	Gly	Val	Thr 965	Leu	Val	Gly	His	Gln 970	Lys	Lуз	Ile	Met	Asn 975	Ser
Leu	Gln	Glu	Met 980	Lуз	Val	Gln	Leu	Val 985	Asn	Gly	Met	Val	Pro 990	Leu	

(2) INFORMATION FOR SEQ ID NO:14:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 3116 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:

 - (A) NAME/KEY: CDS
 (B) LOCATION: 34..2994
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:
- AAGCGGCAGG AGCAGCGTTG GCACCGGCGA ACC ATG GCT GGG ATT TTC TAT TTC 54 Met Ala Gly Ile Phe Tyr Phe 1
- GCC CTA TTT TCG TGT CTC TTC GGG ATT TGC GAC GCT GTC ACA GGT TCC 102 Ala Leu Phe Ser Cys Leu Phe Gly Ile Cys Asp Ala Val Thr Gly Ser 10

															GTT Val	150
						Ile									GAG Glu 55	198
															CAA Gln	246
GTG Val	TGC Cys	AAT Asn	GTG Val 75	ATG Met	GAA Glu	CCC Pro	AGC Ser	CAG Gln 80	AAT Asn	AAC Asn	TGG Trp	CTA Leu	CGA Arg 85	ACT Thr	GAT Asp	294
												GAG Glu 100			TTC Phe	342
ACC Thr	TTG Leu 105	AGG Arg	GAC Asp	TGC Cys	AAT Asn	AGT Ser 110	CTT Leu	CCG Pro	GGC Gly	GTC Val	ATG Met 115	GGG Gly	ACT Thr	TGC Cys	AAG Lys	390
GAG Glu 120	ACG Thr	TTT Phe	AAC Asn	CTG Leu	TAC Tyr 125	TAC Tyr	TAT Tyr	GAA Glu	TCA Ser	GAC Asp 130	AAC Asn	GAC Asp	AAA Lys	GAG Glu	CGT Arg 135	438
TTC Phe	ATC Ile	AGA Arg	GAG Glu	AAC Asn 140	CAG Gln	TTT Phe	GTC Val	AAA Lys	ATT Ile 145	GAC Asp	ACC Thr	ATT Ile	GCT Ala	GCT Ala 150	GAT Asp	486
GAG Glu	AGC Ser	TTC Phe	ACC Thr 155	CAA Gln	GTG Val	GAC Asp	ATT Ile	GGT Gly 160	GAC Asp	AGA Arg	ATC Ile	ATG Met	AAG Lys 165	CTG Leu	AAC Asn	534
ACC Thr	GAG Glu	ATC Ile 170	CGG Arg	GAT Asp	GTA Val	GGG Gly	CCA Pro 175	TTA Leu	AGC Ser	AAA Lys	AAG Lys	GGG Gly 180	TTT Phe	TAC Tyr	CTG Leu	582
GCT Ala	TTT Phe 185	CAG Gln	GAT Asp	GTG Val	GGG Gly	GCC Ala 190	TGC Cys	ATC Ile	GCC Ala	CTG Leu	GTA Val 195	TCA Ser	GTC Val	CGT Arg	GTG Val	630
TTC Phe 200	TAT Tyr	AAA Lys	AAG Lys	TGT Cys	CCA Pro 205	CTC Leu	ACA Thr	GTC Val	CGC Arg	AAT Asn 210	CTG Leu	GCC Ala	CAG Gln	TTT Phe	CCT Pro 215	678
GAC Asp	ACC Thr	ATC Ile	ACA Thr	GGG Gly 220	GCT Ala	GAT Asp	ACG Thr	TCT Ser	TCC Ser 225	CTG Leu	GTG Val	GAA Glu	GTT Val	CGA Arg 230	GGC	726
TCC Ser	TGT Cys	GTC Val	AAC Asn 235	AAC Asn	TCA Ser	GAA Glu	GAG Glu	AAA Lys 240	GAT Asp	GTG Val	CCA Pro	AAA Lys	ATG Met 245	TAC Tyr	TGT Cys	774

									•							
														TGC Cys		822
															GGA Gly	870
														CCA Pro		918
CAC His	AGC Ser	TAC Tyr	TCT Ser	GTC Val 300	TGG Trp	GAA Glu	GGA Gly	GCC Ala	ACC Thr 305	TCG Ser	TGC Cys	ACC Thr	TGT Cys	GAC Asp 310	CGA Arg	966
														ACC Thr		1014
														ACA Thr	TCT Ser	1062
GTG Val	AAC Asn 345	TTG Leu	GAA Glu	TGG Trp	AGT Ser	AGC Ser 350	CCT Pro	CAG Gln	AAT Asn	ACA Thr	GGT Gly 355	GGC Gly	CGC Arg	CAG Gln	GAC Asp	1110
						Cys								CCC Pro		1158
														CAG Gln 390		1206
GGC Gly	TTG Leu	AAG Lys	ACC Thr 395	ACC Thr	AAA Lys	GTC Val	TCC Ser	ATC Ile 400	ACT Thr	GAC Asp	CTC Leu	CTA Leu	GCT Ala 405	CAT His	ACC Thr	1254
														TAT Tyr		1302
CCT Pro	AAC Asn 425	CCA Pro	GAC Asp	CAA Gln	TCA Ser	GTT Val 430	TCT Ser	GTC Val	ACT Thr	GTG Val	ACC Thr 435	ACC Thr	AAC Asn	CAA Gln	GCA Ala	1350
GCA Ala 440	CCA Pro	TCA Ser	TCC Ser	ATT Ile	GCT Ala 445	TTG Leu	GTC Val	CAG Gln	GCT Ala	AAA Lys 450	GAA Glu	GTC Val	ACA Thr	AGA Arg	TAC Tyr 455	1398
AGT Ser	GTG Val	GCA Ala	CTG Leu	GCT Ala 460	TGG Trp	CTG Leu	GAA Glu	CCA Pro	GAT Asp 465	CGG Arg	CCC Pro	AAT Asn	GGG Gly	GTA Val 470	ATC Ile	1446

				-		ТАТ Туг									AGC Ser	1494
						GCT Ala									CTG Leu	1542
						GTT Val 510										1590
						GAG Glu										1638
						GAT Asp										1686
						GTG Val										1734
						AGT Ser										1782
						AAT Asn 590										1830
						AAC Asn										1878
						AAG Lys										1926
						GGG Gly										1974
					_	ACT Thr	_	_			. —					2022
						GAG Glu 670										2070
CCG Pro 680	AAC Asn	ATC Ile	ATT Ile	CAG, His	TTG Leu 685	GAA Glu	GGC Gly	GTG Val	GTC Val	ACT Thr 690	AAA Lys	TGT Cys	AAA Lys	CCA Pro	GTA Val 695	2118

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								TTC Phe 710		2166
								ATG Met		2214
								TAT Tyr		2262
_		_						TTG Leu		2310
								GAT Asp		2358
								TGG Trp 790		2406
								GAT Asp		2454
								GAG Glu		2502
								GAG Glu		2550
								CAC His		2598
								AAA Lys 870		2646
			,					AAC Asn		2694
								TTG Leu		2742
								GAT Asp		2790

		GCC Ala														2838
		ACC Thr														2886
		'ATT Ile														2934
		CAG Gln 970														2982
	CCC Pro 985	GTC Val	TGAC	GCCAC	STA (CTGA <i>I</i>	LAATA	AC TO	CAAAI	ACTC:	r TG	AAAT!	ragt			3031
TTA	CCTC	ATC (CATGO	CACTI	T A	ATTG/	AAGA	CTC	GCAC'	TTTT	TTT	ACTTO	CGT (CTTCC	SCCCTC	3091
TGA.	AATT	AAA (CAAA	rgaa <i>i</i>	LA AJ	AAAA										3116
(2)	INF	ORMA	поп	FOR	SEQ	ID N	10:15	5 :								
		(i) S	(A) (B)	LEN TYE	IGTH:	RACTE : 986 amino GY: 1	ami aci	ino a id		3						
	(:	ii) N	OLEC	CULE	TYPE	E: pi	otei	n								
	(:	ki) S	EQUE	ENCE	DESC	CRIPT	ON:	SEC	DI	NO:1	15:					
Met 1	Ala	Gly	Ile	Phe 5	Tyr	Phe	Ala	Leu	Phe 10	Ser	Cys	Leu	Phe	Gly 15	Ile	
Cys	Asp	Ala	Val 20	Thr	Gly	Ser	Arg	Val 25	Tyr	Pro	Ala	Asn	Glu 30	Val	Thr	
Leu	Leu	Asp 35	Ser	Arg	Ser	Val	Gln 40	Gly	Glu	Leu	Gly	Trp 45	Ile	Ala	Ser	
Pro	Leu	Glu	Gly	Gly	Trp	Glu	Glu	Val	Ser	Ile	Met	Asp	Glu	Lys	Asn	

Thr Pro Ile Arg Thr Tyr Gln Val Cys Asn Val Met Glu Pro Ser Gln 65 70 75 80

Asn Asn Trp Leu Arg Thr Asp Trp Ile Thr Arg Glu Gly Ala Gln Arg

Gly Val Met Gly Thr Cys Lys Glu Thr Phe Asn Leu Tyr Tyr Tyr Glu 120 Ser Asp Asn Asp Lys Glu Arg Phe Ile Arg Glu Asn Gln Phe Val Lys Ile Asp Thr Ile Ala Ala Asp Glu Ser Phe Thr Gln Val Asp Ile Gly 155 Asp Arg Ile Met Lys Leu Asn Thr Glu Ile Arg Asp Val Gly Pro Leu Ser Lys Lys Gly Phe Tyr Leu Ala Phe Gln Asp Val Gly Ala Cys Ile Ala Leu Val Ser Val Arg Val Phe Tyr Lys Lys Cys Pro Leu Thr Val Arg Asn Leu Ala Gln Phe Pro Asp Thr Ile Thr Gly Ala Asp Thr Ser 215 Ser Leu Val Glu Val Arg Gly Ser Cys Val Asn Asn' Ser Glu Glu Lys Asp Val Pro Lys Met Tyr Cys Gly Ala Asp Gly Glu Trp Leu Val Pro Ile Gly Asn Cys Leu Cys Asn Ala Gly His Glu Glu Arg Ser Gly Glu Cys Gln Ala Cys Lys Ile Gly Tyr Tyr Lys Ala Leu Ser Thr Asp Ala 280 Thr Cys Ala Lys Cys Pro Pro His Ser Tyr Ser Val Trp Glu Gly Ala Thr Ser Cys Thr Cys Asp Arg Gly Phe Phe Arg Ala Asp Asn Asp Ala 315 Ala Ser Met Pro Cys Thr Arg Pro Pro Ser Ala Pro Leu Asn Leu Ile Ser Asn Val Asn Glu Thr Ser Val Asn Leu Glu Trp Ser Ser Pro Gln 345 Asn Thr Gly Gly Arg Gln Asp Ile Ser Tyr Asn Val Val Cys Lys 360 Cys Gly Ala Gly Asp Pro Ser Lys Cys Arg Pro Cys Gly Ser Gly Val His Tyr Thr Pro Gln Gln Asn Gly Leu Lys Thr Thr Lys Val Ser Ile 395 Thr Asp Leu Leu Ala His Thr Asn Tyr Thr Phe Glu Ile Trp Ala Val

Asn Gly Val Ser Lys Tyr Asn Pro Asn Pro Asp Gln Ser Val Ser Val Thr Val Thr Thr Asn Gln Ala Ala Pro Ser Ser Ile Ala Leu Val Gln Ala Lys Glu Val Thr Arg Tyr Ser Val Ala Leu Ala Trp Leu Glu Pro Asp Arg Pro Asn Gly Val Ile Leu Glu Tyr Glu Val Lys Tyr Tyr Glu 470 Lys Asp Gln Asn Glu Arg Ser Tyr Arg Ile Val Arg Thr Ala Ala Arg Asn Thr Asp Ile Lys Gly Leu Asn Pro Leu Thr Ser Tyr Val Phe His Val Arg Ala Arg Thr Ala Ala Gly Tyr Gly Asp Phe Ser Glu Pro Leu Glu Val Thr Thr Asn Thr Val Pro Ser Arg Ile Ile Gly Asp Gly Ala 535 Asn Ser Thr Val Leu Leu Val Ser Val Ser Gly Ser Val Val Leu Val 550 Val Ile Leu Ile Ala Ala Phe Val Ile Ser Arg Arg Arg Ser Lys Tyr Ser Lys Ala Lys Gln Glu Ala Asp Glu Glu Lys His Leu Asn Gln Gly Val Arg Thr Tyr Val Asp Pro Phe Thr Tyr Glu Asp Pro Asn Gln Ala Val Arg Glu Phe Ala Lys Glu Ile Asp Ala Ser Cys Ile Lys Ile Glu 615 Lys Val Ile Gly Val Gly Glu Phe Gly Glu Val Cys Ser Gly Arg Leu Lys Val Pro Gly Lys Arg Glu Ile Cys Val Ala Ile Lys-Thr Leu Lys Ala Gly Tyr Thr Asp Lys Gln Arg Arg Asp Phe Leu Ser Glu Ala Ser Ile Met Gly Gln Phe Asp His Pro Asn Ile Ile His Leu Glu Gly Val Val Thr Lys Cys Lys Pro Val Met Ile Ile Thr Glu Tyr Met Glu Asn 695 Gly Ser Leu Asp Ala Phe Leu Arg Lys Asn Asp Gly Arg Phe Thr Val

Ile Gln Leu Val Gly Met Leu Arg Gly Ile Gly Ser Gly Met Lys Tyr Leu Ser Asp Met Ser Tyr Val His Arg Asp Leu Ala Ala Arg Asn Ile Leu Val Asn Ser Asn Leu Val Cys Lys Val Ser Asp Phe Gly Met Ser 760 Arg Val Leu Glu Asp Asp Pro Glu Ala Ala Tyr Thr Thr Arg Gly Gly Lys Ile Pro Ile Arg Trp Thr Ala Pro Glu Ala Ile Ala Tyr Arg Lys 790 795 Phe Thr Ser Ala Ser Asp Val Trp Ser Tyr Gly Ile Val Met Trp Glu Val Met Ser Tyr Gly Glu Arg Pro Tyr Trp Asp Met Ser Asn Gln Asp Val Ile Lys Ala Ile Glu Glu Gly Tyr Arg Leu Pro Pro Met Asp 840 Cys Pro Ile Ala Leu His Gln Leu Met Leu Asp Cys Trp Gln Lys Glu Arg Ser Asp Arg Pro Lys Phe Gly Gln Ile Val Asn Met Leu Asp Lys Leu Ile Arg Asn Pro Asn Ser Leu Lys Arg Thr Gly Thr Glu Ser Ser Arg Pro Asn Thr Ala Leu Leu Asp Pro Ser Ser Pro Glu Phe Ser Ala 900 905 Val Val Ser Val Gly Asp Trp Leu Gln Ala Ile Lys Met Asp Arg Tyr 920 Lys Asp Asn Phe Thr Ala Ala Gly Tyr Thr Thr Leu Glu Ala Val Val His Val Asn Gln Glu Asp Leu Ala Arg Ile Gly Ile Thr Ala Ile Thr His Gln Asn Lys Ile Leu Ser Ser Val Gln Ala Met Arg Thr Gln Met Gln Gln Met His Gly Arg Met Val Pro Val

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(2) IN	FORMATION	FOR	SEQ	ID	NO:16:
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(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 4529 base pairs
(B) TYPE: nucleic acid
(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 186..3182

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

CGGTGCGAGC GAACAGGA	AGT GGGGGGAAA	TTAAAAAAAG CTAA	ACGTGG AGCAGCCGAT	60
CGGGGACCGA GAAGGGGA	AAT CGATGCAAGG	AGCACACTAA AACA	AAAAGCT ACTTCGGAAC	120
AAACAGCATT TAAAAATO	CCA CGACTCAAGA	TAACTGAAAC CTAA	AAATAAA ACCTGCTCAT	180
GCACC ATG GTT TTT (Met Val Phe (C CCT TCA TGG AT r Pro Ser Trp II 10	- · · · · - 	227
TAC ATC TGG CTG CTC Tyr Ile Trp Leu Leu 15				275
AAG GAA GTA CTA CTC Lys Glu Val Leu Leu 35	Leu Asp Ser 1			323
TGG ATT TCC TCT CCA Trp Ile Ser Ser Pro 50				371
GAG AAC TAT ACC CCC Glu Asn Tyr Thr Pro 65				419
CCC AAC CAA AAC AAC Pro Asn Gln Asn Asn 80				467
GCA CAA AGG ATT TTT Ala Gln Arg Ile Phe 95				515
AGT CTT CCT GGA GTA Ser Leu Pro Gly Val 115	Leu Gly Thr (,	563

			GAC Asp						611
			ATT Ile						659
			ATG Met 165						7 07
			GGA Gly						755
			TCT Ser						803
			GCT Ala					_	851
			GAG Glu						899
			GCC Ala 245						947
			GGA Gly						995
			GAA Glu		Arg				1043
			TGC Cys						1091
			AGA Arg						1139
	-		TAC Tyr 325						1187
			AAC Asn						1235

							TAC Tyr 365	AGA Arg	1283
 							CCC		1331
							GAT Asp		1379
							TTT Phe		1427
							AGG Arg		1475
							CAA Gln 445		1523
							CTT Leu		1571
							GAA Glu		1619
							GTA Val		1667
							ACA Thr		1715
							AAT Asn 525		1763
							AAA Lys		1811
							ATC Ile		1859
							GTC Val	TTT Phe	1907

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GGC Gly 575	TTC Phe	ATC Ile	ATT Ile	GGG Gly	AGA Arg 580	AGG Arg	CAC His	TGT Cys	GGT Gly	TAT Tyr 585	AGC Ser	AAA Lys	GCT Ala	GAC Asp	CAA Gln 590	1955
GAA Glu	GGC Gly	GAT Asp	GAA Glu	GAG Glu 595	CTT Leu	TAC Tyr	TTT Phe	CAT His	TTT Phe 600	AAA Lys	TTT Phe	CCA Pro	GGC Gly	ACC Thr 605	AAA Lys	2003
					GAA Glu											2051
					CTA Leu											2099
					TTC Phe											2147
CCA Pro 655	GGG Gly	AAA Lys	AGA Arg	GAT Asp	GTT Val 660	GCA Ala	GTA Val	GCC Ala	ATA Ile	AAA Lys 665	ACC Thr	CTG Leu	AAA Lys	GTT Val	GGT Gly 670	2195
TAC Tyr	ACA Thr	GAA Glu	AAA Lys	CAA Gln 675	AGG Arg	AGA Arg	GAC Asp	TTT Phe	TTG Leu 680	TGT Cys	GAA Glu	GCA Ala	AGC Ser	ATC Ile 685	ATG Met	2243
					CCA Pro											2291
AGA Arg	GGG Gly	AAA Lys 705	CCA Pro	GTC Val	ATG Met	ATA Ile	GTA Val 710	ATA Ile	GAG Glu	TTC Phe	ATG Met	GAA Glu 715	AAT Asn	GGA Gly	GCC Ala	2339
					AGG Arg											2387
TTA Leu 735	GTA Val	GGA Gly	ATG Met	CTG Leu	AGA Arg 740	GGA Gly	ATT Ile	GCT Ala	GCT Ala	GGA Gly 745	ATG Met	AGA Arg	TAT Tyr	TTG Leu	GCT Ala 750	2435
GAT Asp	ATG Met	GGA Gly	TAT Tyr	GTT Val 755	CAC His	AGG Arg	GAC Asp	CTT Leu	GCA Ala 760	GCT Ala	CGC Arg	AAT Asn	ATT Ile	CTT Leu 765	GTC Val	2483
AAC Asn	AGC Ser	AAT Asn	CTC Leu 770	GTT Val	TGT Cys	AAA Lys	GTG Val	TCA Ser 775	GAT Asp	TTT Phe	GGC Gly	CTG Leu	TCC Ser 780	CGA Arg	GTT Val	2531
ATA Ile	GAG Glu	GAT Asp 785	GAT Asp	CCA Pro	GAA Glu	GCT Ala	GTC Val 790	TAT Tyr	ACA Thr	ACT Thr	ACT Thr	GGT Gly 795	GGA Gly	AAA Lys	ATT Ile	2579

		AGG Arg													ACA Thr	2627
		AGT Ser														2675
		GGA Gly														2723
		ATA Ile														2771
		CTT Leu 865														2819
		CCA Pro														2867
		CCA Pro														2915
		CCT Pro														2963
		GGA Gly														3011
		ACG Thr 945														3059
		GAG Glu														3107
		ATC Ile														3155
TTA Leu								TGAI	ATGC	AT I	TCTC	CCTI	T TA	AGGG	GAGAT	3209
TACA	GACI	GC A	AGAG	AACA	G TA	CTGG	CCTI	CAG	TATA	TGC	ATAG	AATG	CT G	CTAG	AAGAC	3269
AAGT	GATO	TC C	TGGG	TCCI	T CC	AACA	GTGA	AGA	GAAG	TTA	TAAG	AAGC	AC C	TATA	GACTT	3329
GAACTCCTAA GTGCCACCAG AATATATAAA AAGGGAATTT AGGATCCACC ATCGGTGGCC												3389				

AGGAAAATAG	CAGTGACAAT	AAACAAAGTA	CTACCTGAAA	AACATCCAAA	CACCTTGAGC	3449
TCTCTAACCT	CCTTTTTGTC	TTATAGACTT	TTTAAAATGT	ACATAAAGAA	TTTAAGAAAG	3509
AATATATTTG	TCAAATAAAA	TCATGATCTT	ATTGTTAAAA	TTAATGAAAT	ATTTTCCTTA	3569
AATATGTGAT	TTCAGACTAT	TCCTTTTTAA	AATCATTTGT	GTTTATTCTT	CATAAGGACT	3629
TTGTTTTAGA	AAGCTGTTTA	TAGCTTTGGA	CCTTTTTAGT	GTTAAATCTG	TAACATTACT	3689
ACACTGGGTA	CCTTTGAAAG	AATCTCAAAT	TTCAAAAGAA	ATAGCATGAT	TGAAGATACA	3749
TCTCTGTTAG	AACATTGGTA	TCCTTTTTGT	GCCATTTTAT	TCTGTTTAAT	CAGTGCTGTT	3809
TTGATATTGT	TTGCTAATTG	GCAGGTAGTC	AAGAAAATGC	AAGTTGCCAA	GAGCTCTGAT	38,69
ATTTTTTAAA	AAGAATTTTT	TTGTAAAGAT	CAGACAACAC	ACTATCTTTT	CAATGAAAA	3929
AGCAATAATG	ATCCATACAT	ACTATAAGGC	ACTTTTAACA	GATTGTTTAT	AGAGTGATTT	3989
TACTAGAAAG	AATTTAATAA	ACTCGAAGTT	TAGGTTTATG	AGTATATAAA	CAAATGAGGC	4049
ACTTCATCTG	AAGAATGTTG	GTGAAGGCAA	GTCTCTGAAA	GCAGAACTAT	CCAGTGTTAT	4109
СТАААААТТА	ATCTGAGCAC	ATCAAGATTT	TTTCATTCTC	GTGACATTAG	GAAATTTAGG	4169
ATAAATAGTT	GACATATATT	TTATATCCTC	TTCTGTTGAA	TGCAGTCCAA	ACATGAAAGG	4229
AAATAATTGT	TTTATATTAT	AACTCTGAAG	CATGATAAAG	GGGCAGTTCA	CAATTTTCAC	4289
CATTTAAACA	CAAATTTGCT	GCACAGAATA	TCACCATTGC	AGTTCAAAAC	AAAACAAAAC	4349
AAAAAGTCTT	TTGTTTGTGA	ACACTGATGC	AAGAAACTTG	TTAAATGAAA	GGACTCTTTA	4409
CCCTAGAAGG	AAGAGGTGAA	GGATCTGGCT	TGTTTTTAAA	GCTTTATTTA	TTAAACCATA	4469
TTATTTGATT	ACTGTGTTAG	AATTTCATAA	GCAATAATTA	AATGTGTCTT	TATGGAATTC	4529

(2) INFORMATION FOR SEQ ID NO:17:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 998 amino acids
 - (B) TYPE: amino acid (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

Met Val Phe Gln Thr Arg Tyr Pro Ser Trp Ile Ile Leu Cys Tyr Ile

Trp Leu Leu Arg Phe Ala His Thr Gly Glu Ala Gln Ala Ala Lys Glu 20

Val Leu Leu Asp Ser Lys Ala Gln Gln Thr Glu Leu Glu Trp Ile Ser Ser Pro Pro Asn Gly Trp Glu Glu Ile Ser Gly Leu Asp Glu Asn Tyr Thr Pro Ile Arg Thr Tyr Gln Val Cys Gln Val Met Glu Pro Asn Gln Asn Asn Trp Leu Arg Thr Asn Trp Ile Ser Lys Gly Asn Ala Gln Arg Ile Phe Val Glu Leu Lys Phe Thr Leu Arg Asp Cys Asn Ser Leu Pro Gly Val Leu Gly Thr Cys Lys Glu Thr Phe Asn Leu Tyr Tyr 120 Glu Thr Asp Tyr Asp Thr Gly Arg Asn Ile Arg Glu Asn Leu Tyr Val Lys Ile Asp Thr Ile Ala Ala Asp Glu Ser Phe Thr Gln Gly Asp Leu Gly Glu Arg Lys Met Lys Leu Asn Thr Glu Val Arg Glu Ile Gly Pro 165 170 Leu Ser Lys Lys Gly Phe Tyr Leu Ala Phe Gln Asp Val Gly Ala Cys Ile Ala Leu Val Ser Val Lys Val Tyr Tyr Lys Lys Cys Trp Ser Ile 200 Ile Glu Asn Leu Ala Ile Phe Pro Asp Thr Val Thr Gly Ser Glu Phe 210 Ser Ser Leu Val Glu Val Arg Gly Thr Cys Val Ser Ser Ala Glu Glu 230 235 Glu Ala Glu Asn Ala Pro Arg Met His Cys Ser Ala Glu Gly Glu Trp Leu Val Pro Ile Gly Lys Cys Ile Cys Lys Ala Gly Tyr Gln Gln Lys Gly Asp Thr Cys Glu Pro Cys Gly Arg Gly Phe Tyr Lys Ser Ser Ser Gln Asp Leu Gln Cys Ser Arg Cys Pro Thr His Ser Phe Ser Asp Lys Glu Gly Ser Ser Arg Cys Glu Cys Glu Asp Gly Tyr Tyr Arg Ala Pro 315 Ser Asp Pro Pro Tyr Val Ala Cys Thr Arg Pro Pro Ser Ala Pro Gln

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Asn Leu Ile Phe Asn Ile Asn Gln Thr Thr Val Ser Leu Glu Trp Ser 345 Pro Pro Ala Asp Asn Gly Gly Arg Asn Asp Val Thr Tyr Arg Ile Leu 360 365 Cys Lys Arg Cys Ser Trp Glu Gln Gly Glu Cys Val Pro Cys Gly Ser Asn Ile Gly Tyr Met Pro Gln Gln Thr Gly Leu Glu Asp Asn Tyr Val 395 Thr Val Met Asp Leu Leu Ala His Ala Asn Tyr Thr Phe Glu Val Glu Ala Val Asn Gly Val Ser Asp Leu Ser Arg Ser Gln Arg Leu Phe Ala Ala Val Ser Ile Thr Thr Gly Gln Ala Ala Pro Ser Gln Val Ser Gly 440 Val Met Lys Glu Arg Val Leu Gln Arg Ser Val Glu Leu Ser Trp Gln Glu Pro Glu His Pro Asn Gly Val Ile Thr Glu Tyr Glu Ile Lys Tyr 470 Tyr Glu Lys Asp Gln Arg Glu Arg Thr Tyr Ser Thr Val Lys Thr Lys Ser Thr Ser Ala Ser Ile Asn Asn Leu Lys Pro Gly Thr Val Tyr Val Phe Gln Ile Arg Ala Phe Thr Ala Ala Gly Tyr Gly Asn Tyr Ser Pro 520 Arg Leu Asp Val Ala Thr Leu Glu Glu Ala Thr Gly Lys Met Phe Glu 535 Ala Thr Ala Val Ser Ser Glu Gln Asn Pro Val Ile Ile Ala Val 555 Val Ala Val Ala Gly Thr Ile Ile Leu Val Phe Met Val Phe Gly Phe Ile Ile Gly Arg Arg His Cys Gly Tyr Ser Lys Ala Asp Gln Glu Gly 585 Asp Glu Glu Leu Tyr Phe His Phe Lys Phe Pro Gly Thr Lys Thr Tyr 600 Ile Asp Pro Glu Thr Tyr Glu Asp Pro Asn Arg Ala Val His Gln Phe 615 Ala Lys Glu Leu Asp Ala Ser Cys Ile Lys Ile Glu Arg Val Ile Gly

Ala Gly Glu Phe Gly Glu Val Cys Ser Gly Arg Leu Lys Leu Pro Gly 650 Lys Arg Asp Val Ala Val Ala Ile Lys Thr Leu Lys Val Gly Tyr Thr Glu Lys Gln Arg Arg Asp Phe Leu Cys Glu Ala Ser Ile Met Gly Gln Phe Asp His Pro Asn Val Val His Leu Glu Gly Val Val Thr Arg Gly 695 Lys Pro Val Met Ile Val Ile Glu Phe Met Glu Asn Gly Ala Leu Asp Ala Phe Leu Arg Lys His Asp Gly Gln Phe Thr Val Ile Gln Leu Val Gly Met Leu Arg Gly Ile Ala Ala Gly Met Arg Tyr Leu Ala Asp Met Gly Tyr Val His Arg Asp Leu Ala Ala Arg Asn Ile Leu Val Asn Ser Asn Leu Val Cys Lys Val Ser Asp Phe Gly Leu Ser Arg Val Ile Glu Asp Asp Pro Glu Ala Val Tyr Thr Thr Thr Gly Gly Lys Ile Pro Val Arg Trp Thr Ala Pro Glu Ala Ile Gln Tyr Arg Lys Phe Thr Ser Ala Ser Asp Val Trp Ser Tyr Gly Ile Val Met Trp Glu Val Met Ser Tyr Gly Glu Arg Pro Tyr Trp Asp Met Ser Asn Gln Asp Val Ile Lys Ala 840 Ile Glu Glu Gly Tyr Arg Leu Pro Ala Pro Met Asp Cys Pro Ala Gly Leu His Gln Leu Met Leu Asp Cys Trp Gln Lys Glu Arg Ala Glu Arg 870 875 Pro Lys Phe Glu Gln Ile Val Gly Ile Leu Asp Lys Met Ile Arg Asn 890 Pro Asn Ser Leu Lys Thr Pro Leu Gly Thr Cys Ser Arg Pro Ile Ser Pro Leu Leu Asp Gln Asn Thr Pro Asp Phe Thr Thr Phe Cys Ser Val 920 Gly Glu Trp Leu Gln Ala Ile Lys Met Glu Arg Tyr Lys Asp Asn Phe 935

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WO 95/28484

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Thr Ala Ala Gly Tyr Asn Ser Leu Glu Ser Val Ala Arg Met Thr Ile

Glu Asp Val Met Ser Leu Gly Ile Thr Leu Val Gly His Gln Lys Lys

Ile Met Ser Ser Ile Gln Thr Met Arg Ala Gln Met Leu His Leu His

Gly Thr Gly Ile Gln Val 995

(2) INFORMATION FOR SEQ ID NO:18:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 976 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

Met Glu Leu Gln Ala Ala Arg Ala Cys Phe Ala Leu Leu Trp Gly Cys

Ala Leu Ala Ala Ala Ala Ala Gln Gly Lys Glu Val Val Leu Leu

Asp Phe Ala Ala Ala Gly Gly Glu Leu Gly Trp Leu Thr His Pro Tyr

Gly Lys Gly Trp Asp Leu Met Gln Asn Ile Met Asn Asp Met Pro Ile

Tyr Met Tyr Ser Val Cys Asn Val Met Ser Gly Asp Gln Asp Asn Trp 65 70 75 80

Leu Arg Thr Asn Trp Val Tyr Arg Gly Glu Ala Glu Arg Asn Asn Phe

Glu Leu Asn Phe Thr Val Arg Asp Cys Asn Ser Phe Pro Gly Gly Ala 105

Ser Ser Cys Lys Glu Thr Phe Asn Leu Tyr Tyr Ala Glu Ser Asp Leu

Asp Tyr Gly Thr Asn Phe Gln Lys Arg Leu Phe Thr Lys Ile Asp Thr 135

Ile Ala Pro Asp Glu Ile Thr Val Ser Ser Asp Phe Glu Ala Arg His

Val	Lys	Leu	Asn	Val 165		Glu	Arg	Ser	Val 170		Pro	Leu	Thr	Arg 175	Lys
Gly	Phe	Tyr	Leu 180	Ala	Phe	Gln	Asp	Ile 185		Ala	Суз	Val	Ala 190		Leu
Ser	Val	Arg 195	Val	Tyr	Tyr	Lys	Lys 200	Cys	Pro	Glu	Leu	Leu 205		Gly	Leu
Ala	His 210		Pro	Glu	Thr	Ile 215	Ala	Gly	Ser	Asp	Ala 220	Pro	Ser	Leu	Ala
Thr 225	Val	Ala	Gly	Thr	Cys 230	Val	Asp	His	Ala	Val 235	Val	Pro	Pro	Gly	Gly 240
Glu	Glu	Pro	Arg	Met 245	His	Суз	Ala	Val	Asp 250	Gly	Glu	Trp	Leu	Val 255	Pro
Ile	Gly	Gln	Cys 260	Leu	Cys	Gln	Ala	Gly 265	Tyr	Glu	Lys	Val	Glu 270	Asp	Ala
Cys	Gln	Ala 275	Суз	Ser	Pro	Gly	Phe 280	Phe	Lys	Phe	Glu	Ala 285	Ser	Glu	Ser
Pro	Cys 290	Leu	Glu	Суз	Pro	Glu 295	His	Thr	Leu	Pro	Ser 300	Pro	Glu	Gly	Ala
Thr 305	Ser	Суѕ	Glu	Cys	Glu 310	Glu	Gly	Phe	Phe	Arg 315	Ala	Pro	Gln	Asp	Pro 320
Ala	Ser	Met		Cys 325	Thr	Arg	Pro	Pro	Ser 330	Ala	Pro	His	Tyr	Leu 335	Thr
Ala	Val	Gly	Met 340	Gly	Ala	Lys	Val	Glu 345	Leu	Arg	Trp	Thr	Pro 350	Pro	Gln
Asp	Ser	Gly 355	Gly	Arg	Glu	Asp	11e 360	Val	Tyr	Ser	Val	Thr 365	Суз	Glu	Gln
Cys	Trp 370	Pro	Glu	Ser	Gly	Glu 375	Cys	Gly	Pro	Суз	Glu 380	Ala	Ser	Val	Arg
Tyr 385	Ser	Glu	Pro	Pro	His 390	Gly	Leu	Thr	Arg	Thr 395	Ser	Val	Thr	Val	Ser 400
Asp	Leu	Glu	Pro	His 405	Met	Asn	Tyr	Thr	Phe 410	Thr	Val	Glu	Ala	Arg 415	Asn
Gly	Val	Ser	Gly 420	Leu	Val	Thr	Ser	Arg 425	Ser	Phe	Arg	Thr	Ala 430	Ser	Val
Ser	Ile	Asn 435	Gln	Thr	Glu	Pro	Pro 440	Lys	Val	Arg	Leu	Glu 445	Gly	Arg	Ser
	Thr	Ser	Leu	Ser		Ser 455	Trp	Ser	Ile		Pro	Pro	Gln	Gln	Ser

Arg 465	Val	Trp	Lys	Tyr	Glu 470	Val	Thr	Tyr	Arg	Lys 475		Gly	Asp	Ser	Ası 480
Ser	Tyr	Asn	Val	Arg 485	Ārg	Thr	Glu	Gly	Phe 490		Val	Thr	Leu	Asp 495	_
Leu	Ala	Pro	Asp 500	Thr	Thr	Tyr	Leu	Val 505	Gln	Val	Gln	Ala	Leu 510	Thr	Glr
Glu	Gly	Gln 515	Gly	Ala	Gly	Ser	Lys 520	Val	His	Glu	Phe	Gln 525	Thr	Leu	Sei
Pro	Glu 530	Gly	Ser	Gly	Asn	Leu 535	Ala	Val	Ile	Gly	Gly 540	Val	Ala	Val	Gl
Val 545	Val	Leu	Leu	Leu	Val 550	Leu	Ala	Gly	Val	Gly 555	Phe	Phe	Ile	His	Arg 560
Arg	Arg	Lys	Asn	Gln 565	Arg	Ala	Arg	Gln	Ser 570	Pro	Glu	Asp	Val	Tyr 575	Phe
Ser	Lys	Ser	Glu 580	Gln	Leu	Lys	Pro	Leu 585	Lys	Thr	Tyr	Val	Asp 590	Pro	His
		Glu 595					600					605			
	610	Ser				615					620				
625		Val			630					635					640
		Val		645					650					655	
		Asp	660					665					670		
		Ile 675					680					685			
	690	Ile			*	695					700				
705		Lys			710					715					720
		Ile		725					730					735	
٠.		Asp	740					745					750		
Cys	Lys	Val 755	Ser	Asp	Phe	Gly	Leu 760	Ser	Arg	Val		Glu 765	Asp	Asp	Pro

- Glu Ala Thr Tyr Thr Thr Ser Gly Gly Lys Ile Pro Ile Arg Trp Thr Ala Pro Glu Ala Ile Ser Tyr Arg Lys Phe Thr Ser Ala Ser Asp Val 790 Trp Ser Phe Gly Ile Val Met Trp Glu Val Met Thr Tyr Gly Glu Arg Pro Tyr Trp Glu Leu Ser Asn His Glu Val Met Lys Ala Ile Asn Asp 825 Gly Phe Arg Leu Pro Thr Pro Met Asp Cys Pro Ser Ala Ile Tyr Gln Leu Met Met Gln Cys Trp Gln Gln Glu Arg Ala Arg Arg Pro Lys Phe Ala Asp Ile Val Ser Ile Leu Asp Lys Leu Ile Arg Ala Pro Asp Ser 870 Leu Lys Thr Leu Ala Asp Phe Asp Pro Arg Val Ser Ile Arg Leu Pro 885 890 Ser Thr Ser Gly Ser Glu Gly Val Pro Phe Arg Thr Val Ser Glu Trp 905 Leu Glu Ser Ile Lys Met Gln Gln Tyr Thr Glu His Phe Met Ala Ala 915 920 Gly Tyr Thr Ala Ile Glu Lys Val Val Gln Met Thr Asn Asp Asp Ile Lys Arg Ile Gly Val Arg Leu Pro Gly His Gln Lys Arg Ile Ala Tyr 950 Ser Leu Leu Gly Leu Lys Asp Gln Val Asn Thr Val Gly Ile Pro Ile
- (2) INFORMATION FOR SEQ ID NO:19:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 984 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:
 - Met Glu Arg Arg Trp Pro Leu Gly Leu Gly Leu Val Leu Leu Cys
 1 5 10 15

Ala Pro Leu Pro Pro Gly Ala Arg Ala Lys Glu Val Thr Leu Met Asp Thr Ser Lys Ala Gln Gly Glu Leu Gly Trp Leu Leu Asp Pro Pro Lys Asp Gly Trp Ser Glu Gln Gln Gln Ile Leu Asn Gly Thr Pro Leu Tyr Met Tyr Gln Asp Cys Pro Met Gln Gly Arg Arg Asp Thr Asp His Trp Leu Arg Ser Asn Trp Ile Tyr Arg Gly Glu Glu Ala Ser Arg Val His Val Glu Leu Gln Phe Thr Val Arg Asp Cys Lys Ser Phe Pro Gly Gly 105 Ala Gly Pro Leu Gly Cys Lys Glu Thr Phe Asn Leu Leu Tyr Met Glu Ser Asp Gln Asp Val Gly Ile Gln Leu Arg Arg Pro Leu Phe Gln Lys Val Thr Thr Val Ala Ala Asp Gln Ser Phe Thr Ile Arg Asp Leu Ala 150 Ser Gly Ser Val Lys Leu Asn Val Glu Arg Cys Ser Leu Gly Arg Leu Thr Arg Arg Gly Leu Tyr Leu Ala Phe His Asn Pro Gly Ala Cys Val 185 Ala Leu Val Ser Val Arg Val Phe Tyr Gln Arg Cys Pro Glu Thr Leu Asn Gly Leu Ala Gln Phe Pro Asp Thr Leu Pro Gly Pro Ala Gly Leu Val Glu Val Ala Gly Thr Cys Leu Pro His Ala Arg Ala Ser Pro Arg Pro Ser Gly Ala Pro Arg Met His Cys Ser Pro Asp Gly Glu Trp Leu Val Pro Val Gly Arg Cys His Cys Glu Pro Gly Tyr Glu Glu Gly Gly 265 Ser Gly Glu Ala Cys Val Ala Cys Pro Ser Gly Ser Tyr Arg Met Asp Met Asp Thr Pro His Cys Leu Thr Cys Pro Gln Gln Ser Thr Ala Glu 295 Ser Glu Gly Ala Thr Ile Cys Thr Cys Glu Ser Gly His Tyr Arg Ala

Pro Gly Glu Gly Pro Gln Val Ala Cys Thr Gly Pro Pro Ser Ala Pro 325 330 Arg Asn Leu Ser Phe Ser Ala Ser Gly Thr Gln Leu Ser Leu Arg Trp Glu Pro Pro Ala Asp Thr Gly Gly Arg Gln Asp Val Arg Tyr Ser Val Arg Cys Ser Gln Cys Gln Gly Thr Ala Gln Asp Gly Gly Pro Cys Gln Pro Cys Gly Val Gly Val His Phe Ser Pro Gly Ala Arg Ala Leu Thr Thr Pro Ala Val His Val Asn Gly Leu Glu Pro Tyr Ala Asn Tyr Thr 410 Phe Asn Val Glu Ala Gln Asn Gly Val Ser Gly Leu Gly Ser Ser Gly His Ala Ser Thr Ser Val Ser Ile Ser Met Gly His Ala Glu Ser Leu 440 Ser Gly Leu Ser Leu Arg Leu Val Lys Lys Glu Pro Arg Gln Leu Glu 455 Leu Thr Trp Ala Gly Ser Arg Pro Arg Ser Pro Gly Ala Asn Leu Thr 470 Tyr Glu Leu His Val Leu Asn Gln Asp Glu Glu Arg Tyr Gln Met Val 490 Leu Glu Pro Arg Val Leu Leu Thr Glu Leu Gln Pro Asp Thr Thr Tyr 505 Ile Val Arg Val Arg Met Leu Thr Pro Leu Gly Pro Gly Pro Phe Ser Pro Asp His Glu Phe Arg Thr Ser Pro Pro Val Ser Arg Gly Leu Thr Gly Gly Glu Ile Val Ala Val Ile Phe Gly Leu Leu Gly Ala Ala 545 Leu Leu Gly Ile Leu Val Phe Arg Ser Arg Arg Ala Gln Arg Gln 570 Arg Gln Gln Arg His Val Thr Ala Pro Pro Met Trp Ile Glu Arg Thr 585 Ser Cys Ala Glu Ala Leu Cys Gly Thr Ser Arg His Thr Arg Thr Leu 600 His Arg Glu Pro Trp Thr Leu Pro Gly Gly Trp Ser Asn Phe Pro Ser

Arg Glu Leu 625 Gly Glu Phe	·	Ala Trp 630	Leu Met		sp Thr	Val Ile	Gly	Glu 640
Gly Glu Phe	Clu Clu							040
	645	Val Tyr	Arg Gly	Thr L	eu Arg	Leu Pro	Ser 655	Gln
Asp Cys Lys	Thr Val	Ala Ile	Lys Thr		ys Asp	Thr Ser 670		Gly
Gly Gln Trp 675	Trp Asn	Phe Leu	Arg Glu 680	Ala T	hr Ile	Met Gly 685	Gln	Phe
Ser His Pro	His Ile	Leu His 695	Leu Glu	Gly V	al Val 700	Thr Lys	Arg	Lys
Pro Ile Met 705	Ile Ile	Thr Glu 710	Phe Met		sn Ala 15	Ala Leu	Asp	Ala 720
Phe Leu Arg	Glu Arg 725	Glu Asp	Gln Leu	Val P. 730	ro Gly	Gln Leu	Val 735	Ala
Met Leu Gln	Gly Ile 740	Ala Ser	Gly Met 745		yr Leu	Ser Asn 750	His	Asn
Tyr Val His	Arg Asp	Leu Ala	Ala Arg 760	Asn I	le Leu	Val Asn 765	Gln	Asn
Leu Cys Cys 770	Lys Val	Ser Asp 775	Phe Gly	Leu T	hr Arg 780	Leu Leu	Asp	Asp
Phe Asp Gly 785	Thr Tyr	Glu Thr 790	Gln Gly		ys Ile 95	Pro Ile	Arg	Trp 800
Thr Ala Pro	Glu Ala 805	Ile Ala	His Arg	Ile P	he Thr	Thr Ala	Ser 815	qsA
Val Trp Ser	Phe Gly 820	Ile Val	Met Trp 825		al Leu	Ser Phe 830	Gly	Asp
Lys Pro Tyr 835	Gly Glu	Met Ser	Asn Gln 840	Glu V	al Met	Lys Ser 845	Ile	Glu
Asp Gly Tyr 850	Arg Leu	Pro Pro 855	Pro Val	Asp C	ys Pro 860	Ala Pro	Leu	Tyr
Glu Leu Met 865	Lys Asn	Cys Trp 870	Ala Tyr		rg Ala 75	Arg Arg	Pro	His 880
Phe Gln Lys	Leu Gln 885	Ala His	Leu Glu	Gln Le 890	eu Leu	Ala Asn	Pro 895	His
Ser Leu Arg	Thr Ile	Ala Asn	Phe Asp		rg Val	Thr Leu 910	Arg	Leu
Pro Ser Leu 915		Ser Asp	Gly Ile 920	Pro T	yr Arg	Thr Val 925	Ser	Glu

Trp Leu Glu Ser Ile Arg Met Lys Arg Tyr Ile Leu His Phe His Ser

Ala Gly Leu Asp Thr Met Glu Cys Val Leu Glu Leu Thr Ala Glu Asp

Leu Thr Gln Met Gly Ile Thr Leu Pro Gly His Gln Lys Arg Ile Leu

Cys Ser Ile Gln Gly Phe Lys Asp 980

(2) INFORMATION FOR SEQ ID NO:20:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 998 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

Met Ala Arg Ala Arg Pro Pro Pro Pro Pro Ser Pro Pro Pro Gly Leu

Leu Pro Leu Leu Pro Pro Leu Leu Leu Pro Leu Leu Leu Pro 25

Ala Gly Cys Arg Ala Leu Glu Glu Thr Leu Met Asp Thr Lys Trp Val 40

Thr Ser Glu Leu Ala Trp Thr Ser His Pro Glu Ser Gly Trp Glu Glu 55

Val Ser Gly Tyr Asp Glu Ala Met Asn Pro Ile Arg Thr Tyr Gln Val

Cys Asn Val Arg Glu Ser Ser Gln Asn Asn Trp Leu Arg Thr Gly Phe 90

Ile Trp Arg Arg Asp Val Gln Arg Val Tyr Val Glu Leu Lys Phe Thr

Val Arg Asp Cys Asn Ser Ile Pro Asn Ile Pro Gly Ser Cys Lys Glu

Thr Phe Asn Leu Phe Tyr Tyr Glu Ala Asp Ser Asp Val Ala Ser Ala

Ser Ser Pro Phe Trp Met Glu Asn Pro Tyr Val Lys Val Asp Thr Ile

Ala Pro Asp Glu Ser Phe Ser Arg Leu Asp Ala Gly Arg Val Asn Thr 170 165

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Lys Val Arg Ser Phe Gly Pro Leu Ser Lys Ala Gly Phe Tyr Leu Ala 185 Phe Gln Asp Gln Gly Ala Cys Met Ser Leu Ile Ser Val Arg Ala Phe 200 Tyr Lys Lys Cys Ala Ser Thr Thr Ala Gly Phe Ala Leu Phe Pro Glu Thr Leu Thr Gly Ala Glu Pro Thr Ser Leu Val Ile Ala Pro Gly Thr 230 Cys Ile Pro Asn Ala Val Glu Val Ser Val Pro Leu Lys Leu Tyr Cys Asn Gly Asp Gly Glu Trp Met Val Pro Val Gly Ala Cys Thr Cys Ala 265 Thr Gly His Glu Pro Ala Ala Lys Glu Ser Gln Cys Arg Pro Cys Pro 280 Pro Gly Ser Tyr Lys Ala Lys Gln Gly Glu Gly Pro Cys Leu Pro Cys Pro Pro Asn Ser Arg Thr Thr Ser Pro Ala Ala Ser Ile Cys Thr Cys His Asn Asn Phe Tyr Arg Ala Asp Ser Asp Ser Ala Asp Ser Ala Cys Thr Thr Val Pro Ser Pro Pro Arg Gly Val Ile Ser Asn Val Asn Glu 345 Thr Ser Leu Ile Leu Glu Trp Ser Glu Pro Arg Asp Leu Gly Val Arg Asp Asp Leu Leu Tyr Asn Val Ile Cys Lys Lys Cys His Gly Ala Gly 375 Gly Ala Ser Ala Cys Ser Arg Cys Asp Asp Asn Val Glu Phe Val Pro Arg Gln Leu Gly Leu Ser Glu Pro Arg Val His Thr Ser His Leu Leu Ala His Thr Arg Tyr Thr Phe Glu Val Gln Ala Val Asn Gly Val Ser Gly Lys Ser Pro Leu Pro Pro Arg Tyr Ala Ala Val Asn Ile Thr Thr Asn Gln Ala Ala Pro Ser Glu Val Pro Thr Leu Arg Leu His Ser Ser Ser Gly Ser Ser Leu Thr Leu Ser Trp Ala Pro Pro Glu Arg Pro Asn 475

Gly	Val	Ile	Leu	Asp 485		Glu	Met	Lys	190 490		Glu	Lys	Ser	Glu 495	Gly
Ile	Ala	Ser	Thr 500	Val	Thr	Ser	Gln	Met 505		Ser	Val	Gln	Leu 510		Gly
Leu	Arg	Pro 515	Asp	Ala	Arg	Tyr	Val 520		Gln	Val	Arg	Ala 525		Thr	Val
Ala	Gly 530		Gly	Gln	Tyr	Ser 535		Pro	Ala	Glu	Phe 540	Glu	Thr	Thr	Ser
Glu 545	Arg	Gly	Ser	Gly	Ala 550	Gln	Gln	Leu	Gln	Glu 555	Gln	Leu	Pro	Leu	Ile 560
Val	Gly	Ser	Ala	Thr 565	Ala	Gly	Leu	Val	Phe 570	Val	Val	Ala	Val	Val 575	Val
Ile	Ala	Ile	Val 580	Cys	Leu	Arg	Lys	Gln 585	Arg	His	Gly	Ser	Asp 590	Ser	Glu
		595					600					605		Val	_
	610					615					620				Phe
625					630					635				Ile	640
				645					650					655	Gly
			660					665					670	Tyr	
		675					680					685		Gly	
	690					695					700			Lys	
705					710					715				Leu	720
				725					730					Leu 735	
			740					745					750	Glu	
		755					760					765		Asn	
Asn	Leu 770	Val	Cys	Lys	Val	Ser 7 75	Asp	Phe	Gly		Ser 780	Arg	Phe	Leu	Glu

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Asp Asp Pro Ser Asp Pro Thr Tyr Thr Ser Ser Leu Gly Gly Lys Ile 785 790 795 800

Pro Ile Arg Trp Thr Ala Pro Glu Ala Ile Ala Tyr Arg Lys Phe Thr 805 810 815

Ser Ala Ser Asp Val Trp Ser Tyr Gly Ile Val Met Trp Glu Val Met 820 825 830

Ser Tyr Gly Glu Arg Pro Tyr Trp Asp Met Ser Asn Gln Asp Val Ile 835 840 845

Asn Ala Val Glu Gln Asp Tyr Arg Leu Pro Pro Pro Met Asp Cys Pro 850 855 860

Thr Ala Leu His Gln Leu Met Leu Asp Cys Trp Val Arg Asp Arg Asn 865 870 875 880

Leu Arg Pro Lys Phe Ser Gln Ile Val Asn Thr Leu Asp Lys Leu Ile 885 890 895

Arg Asn Ala Ala Ser Leu Lys Val Ile Ala Ser Ala Gln Ser Gly Met 900 905 910

Ser Gln Pro Leu Leu Asp Arg Thr Val Pro Asp Tyr Thr Thr Phe Thr 915 920 925

Thr Val Gly Asp Trp Leu Asp Ala Ile Lys Met Gly Arg Tyr Lys Glu 930 935 940

Ser Phe Val Ser Ala Gly Phe Ala Ser Phe Asp Leu Val Ala Gln Met 945 950 955 960

Thr Ala Glu Asp Leu Leu Arg Ile Gly Val Thr Leu Ala Gly His Gln 965 970 975

Lys Lys Ile Leu Ser Ser Ile Gln Asp Met Arg Leu Gln Met Asn Gln 980 985 990

Thr Leu Pro Val Gln Val 995

(2) INFORMATION FOR SEQ ID NO:21:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 983 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

Met Asp Cys Gln Leu Ser Ile Leu Leu Leu Leu Ser Cys Ser Val Leu 1 5 10 15

Asp Ser Phe Gly Glu Leu Ile Pro Gln Pro Ser Asn Glu Val Asn Leu Leu Asp Ser Lys Thr Ile Gln Gly Glu Leu Gly Trp Ile Ser Tyr Pro Ser His Gly Trp Glu Glu Ile Ser Gly Val Asp Glu His Tyr Thr Pro Ile Arg Thr Tyr Gln Val Cys Asn Val Met Asp His Ser Gln Asn Asn Trp Leu Arg Thr Asn Trp Val Pro Arg Asn Ser Ala Gln Lys Ile Tyr Val Glu Leu Lys Phe Thr Leu Arg Asp Cys Asn Ser Ile Pro Leu Val Leu Gly Thr Cys Lys Glu Thr Phe Asn Leu Tyr Tyr Met Glu Ser Asp 120 Asp Asp His Gly Val Lys Phe Arg Glu His Gln Phe Thr Lys Ile Asp 135 Thr Ile Ala Ala Asp Glu Ser Phe Thr Gln Met Asp Leu Gly Asp Arg Ile Leu Lys Leu Asn Thr Glu Ile Arg Glu Val Gly Pro Val Asn Lys 170 Lys Gly Phe Tyr Leu Ala Phe Gln Asp Val Gly Ala Cys Val Ala Leu Val Ser Val Arg Val Tyr Phe Lys Lys Cys Pro Phe Thr Val Lys Asn 200 Leu Ala Met Phe Pro Asp Thr Val Pro Met Asp Ser Gln Ser Leu Val Glu Val Arg Gly Ser Cys Val Asn Asn Ser Lys Glu Glu Asp Pro Pro 230 Arg Met Tyr Cys Ser Thr Glu Gly Glu Trp Leu Val Pro Ile Gly Lys Cys Ser Cys Asn Ala Gly Tyr Glu Glu Arg Gly Phe Met Cys Gln Ala Cys Arg'Pro Gly Phe Tyr Lys Ala Leu Asp Gly Asn Met Lys Cys Ala Lys Cys Pro Pro His Ser Ser Thr Gln Glu Asp Gly Ser Met Asn Cys Arg Cys Glu Asn Asn Tyr Phe Arg Ala Asp Lys Asp Pro Pro Ser Met 315

Ala Cys Thr Arg Pro Pro Ser Ser Pro Arg Asn Val Ile Ser Asn Ile 330 Asn Glu Thr Ser Val Ile Leu Asp Trp Ser Trp Pro Leu Asp Thr Gly Gly Arg Lys Asp Val Thr Phe Asn Ile Ile Cys Lys Lys Cys Gly Trp 360 Asn Ile Lys Gln Cys Glu Pro Cys Ser Pro Asn Val Arg Phe Leu Pro 375 Arg Gln Phe Gly Leu Thr Asn Thr Thr Val Thr Val Thr Asp Leu Leu Ala His Thr Asn Tyr Thr Phe Glu Ile Asp Ala Val Asn Gly Val Ser Glu Leu Ser Ser Pro Pro Arg Gln Phe Ala Ala Val Ser Ile Thr Thr Asn Gln Ala Ala Pro Ser Pro Val Leu Thr Ile Lys Lys Asp Arg Thr Ser Arg Asn Ser Ile Ser Leu Ser Trp Gln Glu Pro Glu His Pro Asn 455 Gly Ile Ile Leu Asp Tyr Glu Val Lys Tyr Tyr Glu Lys Gln Glu Gln Glu Thr Ser Tyr Thr Ile Leu Arg Ala Arg Gly Thr Asn Val Thr Ile Ser Ser Leu Lys Pro Asp Thr Ile Tyr Val Leu Gln Ile Arg Ala Arg Thr Ala Ala Gly Tyr Gly Thr Asn Ser Arg Lys Phe Glu Phe Glu Thr Ser Pro Asp Ser Phe Ser Ile Ser Gly Glu Ser Ser Gln Val Val Met 535 Ile Ala Ile Ser Ala Ala Val Ala Ile Ile Leu Leu Thr Val Val Ile 550 Tyr Val Leu Ile Gly Arg Phe Cys Gly Tyr Lys Ser Lys His Gly Ala Asp Glu Lys Arg Leu His Phe Gly Asn Gly His Leu Lys Leu Pro Gly Leu Arg Thr Tyr Val Asp Pro His Thr Tyr Glu Asp Pro Thr Gln Ala 600 Val His Glu Phe Ala Lys Glu Leu Asp Ala Thr Asn Ile Ser Ile Asp 610 615

Lys Val Val Gly Ala Gly Glu Phe Gly Glu Val Cys Ser Gly Arg Leu Lys Leu Pro Ser Lys Lys Glu Ile Ser Val Ala Ile Lys Thr Leu Lys Val Gly Tyr Thr Glu Lys Gln Arg Arg Asp Phe Leu Gly Glu Ala Ser 665 Ile Met Gly Gln Phe Asp His Pro Asn Ile Ile Arg Leu Glu Gly Val 680 Val Thr Lys Ser Lys Pro Val Met Ile Val Thr Glu Tyr Met Glu Asn Gly Ser Leu Asp Ser Phe Leu Arg Lys His Asp Ala Gln Phe Thr Val Ile Gln Leu Val Gly Met Leu Arg Gly Ile Ala Ser Gly Met Lys Tyr Leu Ser Asp Met Gly Tyr Val His Arg Asp Leu Ala Ala Arg Asn Ile 745 Leu Ile Asn Ser Asn Leu Val Cys Lys Val Ser Asp Phe Gly Leu Ser Arg Val Leu Glu Asp Asp Pro Glu Ala Ala Tyr Thr Thr Arg Gly Gly Lys Ile Pro Ile Arg Trp Thr Ser Pro Glu Ala Ile Ala Tyr Arg Lys Phe Thr Ser Ala Ser Asp Val Trp Ser Tyr Gly Ile Val Leu Trp Glu Val Met Ser Tyr Gly Glu Arg Pro Tyr Trp Glu Met Ser Asn Gln Asp Val Ile Lys Ala Val Asp Glu Gly Tyr Arg Leu Pro Pro Pro Met Asp Cys Pro Ala Ala Leu Tyr Gln Leu Met Leu Asp Cys Trp Gln Lys Asp 855 Arg Asn Asn Arg Pro Lys Phe Glu Gln Ile Val Ser Ile Leu Asp Lys Leu Ile Arg Asn Pro Gly Ser Leu Lys Ile Ile Thr Ser Ala Ala Ala Arg Pro Ser Asn Leu Leu Leu Asp Gln Ser Asn Val Asp Ile Ser Thr 905 Phe Arg Thr Thr Gly Asp Trp Leu Asn Gly Val Arg Thr Ala His Cys 915 920

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	Lys	Glu 930	Ile	Phe	Thr	Gly	Val 935	Glu	Tyr	Ser	Ser	Cys 940	Asp	Thr	Île	Ala	
	Lys 945	Ile	Ser	Thr	Asp	Asp 950	Met	Lys	Lys	Val	Gly 955	Val	Thr	Val	Val	Gly 960	
	Pro	Gln	Lуз	Lys	Ile 965	Ile	Ser	Ser	Ile	Lys 970	Ala	Leu	Glu	Thr	Gln 975	Ser	
	Lys	Asn	Gly	Pro 980	Val	Pro	Val										
(2)	INFO	RMATI	ON E	FOR S	SEQ :	ID NO	22:02	:									
	(i)	(B)	LEN TYI STI	NGTH PE: 1 RANDI	ARACT 24 nucle EDNES	base eic a	e par acid sing	irs									
	(ii)	MOLE	CULE	E TYI	PE: o	DNA											
	(xi)	SEQU	JENCI	E DES	SCRII	PTIO	N: SI	EQ II	ON C	:22:		,					
CTG	CTCGC	cg co	GTG	GAAG	AA A	CG											24
(2)	INFO	RMATI	ON I	FOR S	SEQ :	ID N	0:23	:									
	(i)	(B)	LEI TYI STI	NGTH PE: 1 RANDI	ARACT : 39 nucle EDNES	base eic a SS: :	e par acid sing	irs									
	(ii)	MOLE	CUL	E TY	PE: 0	DNA						,					
	(xi)	SEQU	JENCI	E DES	SCRII	PTIO	N: SI	EQ II	O NO:	:23:							
GCG	CTAG	AT T	ATCAC	CTTC	r cc	rgga:	rgct	TGT	CTGG:	ra							39
(2)	INFO	RMATI	ON I	FOR S	SEQ :	ID N	0:24	:	•								
	(i)	(B)	LEN TYI STI	NGTH PE: 1 RANDI	ARAC: : 48 nucle EDNES	base eic a	e pa: acid sing:	irs		*							

(ii) MOLECULE TYPE: cDNA

	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:24:	
GCGC	GACGC	CG CCGCCATGGC CCTGGATTGC CTGCTGCTGT TCCTCCTG	48
(2)	INFO	RMATION FOR SEQ ID NO:25:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 54 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: cDNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:25:	
CGTI	TCTT	CC ACGGCGGCGA GCAGAGATGC CAGGAGGAAC AGCAGCAGGC AATC	54
(2)	INFO	RMATION FOR SEQ ID NO:26:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 13 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: protein	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:26:	
	Met 1	Ala Leu Asp Cys Leu Leu Leu Phe Leu Leu Ala Ser 5 10	
(2)	INFO	RMATION FOR SEQ ID NO:27:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
•	(ii)	MOLECULE TYPE: cDNA	

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

AGGGAATTCC AYCGNGAYYT NGCNGC

- (2) INFORMATION FOR SEQ ID NO:28:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 24 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

AGGGGATCCR WARSWCCANA CRTC

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WHAT IS CLAIMED IS:

- 1. An isolated nucleic acid encoding a polypeptide having at least one of the biological activities of an EPH-like receptor protein tyrosine kinase, the nucleic acid selected from the group consisting of:
- (a) the nucleic acids set forth in any of SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14 or SEQ ID NO: 16 and their complementary strands;
- (b) a nucleic acid hybridizing to the coding regions of the nucleic acids in any of SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14 or SEQ ID NO: 16; and
- (c) a nucleic acid of (b) which, but for the degeneracy of the genetic code, would hybridize to the coding regions of the nucleic acids in any of SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14 or SEQ ID NO: 16.
- A polypeptide product of expression of a
 nucleic acid of Claim 1 in a procaryotic or eucaryotic host cell.
 - 3. A nucleic acid of Claim 1 which is of human origin.

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4. A nucleic acid of Claim 1 which encodes a polypeptide having part or all of the amino acid sequence as shown in any of SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14 or SEQ ID NO: 16.

- 5. A nucleic acid of Claim 1 encoding a fragment comprising an EPH-like receptor extracellular domain.
- 35 6. A nucleic acid of Claim 1 which is cDNA, genomic DNA, synthetic DNA or RNA.

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7. A nucleic acid of Claim 1 which includes one or more codons preferred for expression in E. coli host cells.

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- 8. A nucleic acid of Claim 1 which includes one or more codon preferred for expression in mammalian cells.
- 9. A nucleic acid encoding amino acids 6-524 as set forth in SEQ ID NO: 10, and optionally encoding an amino terminal methionyl residue.
- 10. A nucleic acid encoding amino acids 1-547
 15 as set forth in SEQ ID NO: 12, and optionally encoding an amino acid terminal methionyl residue.
- 11. A nucleic acid encoding amino acids 21-547 as set forth in SEQ ID NO: 14, and optionally20 encoding an amino terminal methionyl residue.
 - 12. A nucleic acid encoding amino acids 23-553 as set forth in SEQ ID NO: 16, and optionally encoding an amino terminal methionyl residue.

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13. A nucleic acid encoding a chimeric protein, wherein the protein comprises an EPH-like receptor extracellular domain fused to a heterologous receptor cytoplasmic domain.

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14. A nucleic acid of Claim 13 wherein the extracellular domain is selected from the group consisting of HEK5, HEK7, HEK8 and HEK11 extracellular domains.

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- 15. A biologically functional plasmid or viral DNA vector including a nucleic acid of Claim 1.
- 16. A procaryotic or eucaryotic host cell 5 stably transformed or transfected with the plasmid of Claim 15.
- 17. A method of producing an EPH-like receptor protein tyrosine kinase comprising culturing the host cell of Claim 16 to allow the host cell to express the EPH-like receptor protein tyrosine kinase.
- 18. An isolated polypeptide having an amino acid sequence as shown in any of SEQ ID NO: 10, SEQ ID NO: 12, SEQ ID NO: 14 or SEQ ID NO: 16, or a fragment or analog thereof, wherein the polypeptide has at least one of the biological activities of an EPH-like receptor protein tyrosine kinase.
- 20 19. Purified and isolated HEK5 receptor.

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- 20. Purified and isolated HEK7 receptor.
- 21. Purified and isolated HEK8 receptor.
- 22. Purified and isolated HEKll receptor.
- 23. A polypeptide of Claim 18 wherein the biological activity is the binding of a ligand.
- 24. A polypeptide of Claim 18 which is of human origin.
- 25. A polypeptide of Claims 18 characterized 35 by being the product of procaryotic or eucaryotic expression of an exogenous DNA sequence.

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- 26. A polypeptide of Claim 25 wherein the exogenous DNA is a cDNA.
- 5 27. A polypeptide of Claim 25 wherein the exogenous DNA is a genomic DNA.
 - 28. An antibody or fragment thereof specifically binding a polypeptide of Claim 18.

- 29. An antibody of Claim 28 which is a monoclonal antibody.
- 30. A pharmaceutical composition comprising a therapeutically effective amount of a polypeptide of Claim 18 in a mixture with a pharmaceutically acceptable adjuvant, carrier, solubilizer or diluent.
- 31. A pharmaceutical composition comprising a therapeutically effective amount of an antibody of Claim 28 in a mixture with a pharmaceutically acceptable adjuvant, carrier, solubilizer or diluent.
- 32. A method for modulating the endogenous activation of an EPH-like receptor protein tyrosine kinase comprising administering an effective amount of a polypeptide of Claim 18.
- 33. A method for modulating the synthesis of an EPH-like receptor protein tyrosine kinase comprising hybridizing an antisense oligonucleotide to a nucleic acid of Claim 1.

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- 34. A method of identifying a ligand that binds to a receptor polypeptide of Claim 18 comprising the steps of:
- a) exposing at least one molecule to the 5 receptor polypeptide for a time sufficient to allow formation of a receptor/ligand complex;
 - b) removing non-complexed molecules; and
 - c) detecting the presence of the molecule bound to the receptor polypeptide.

1/33 FIG. IA

			-	· 10	5. I	А				
							TCC Ser			48
							GGG Gly			96
							ACG Thr			144
							CGG Arg 60			192
							ATG Met			240
							TCC Ser			288
							TCG Ser			336
							GTG Val			384
							GGC Gly 140			432
							TCC Ser			480
							TCC Ser			528
							CAG Gln			576

2 / 33 FIG. IB

							I		ו .כ							
TTC Phe	CAG Gln	GAA Glu 195	Thr	CTG Leu	TCG Ser	GGG Gly	GCT Ala 200	Glu	AGC Ser	ACA Thr	TCG Ser	CTG Leu 205	Val	GCT Ala	GCC Ala	624
CGG Arg	GGC Gly 210	Ser	TGC Cys	ATC Ile	GCC Ala	AAT Asn 215	GCG Ala	GAA Glu	GAG Glu	GTG Val	GAT Asp 220	GTA Val	CCC Pro	ATC	AAG Lys	672
CTC Leu 225	TAC Tyr	TGT Cys	AAC Asn	GGG Gly	GAC Asp 230	GGC Gly	GAG Glu	TGG Trp	CTG Leu	GTG Val 235	CCC Pro	ATC Ile	GGG Gly	CGC Arg	TGC Cys 240	720
			GCA Ala												CGA Arg	768
GGT Gly	TGT Cys	CCA Pro	TCT Ser 260	GGG Gly	ACT Thr	TTC Phe	AAG Lys	GCC Ala 265	AAC Asn	CAA Gln	GGG Gly	GAT Asp	GAG Glu 270	GCC Ala	TGT Cys	816
			CCC Pro													864
TGT Cys	GTC Val 290	TGC Cys	CGC Arg	AAT Asn	GGC Gly	TAC Tyr 295	TAC Tyr	AGA Arg	GCA Ala	GAC Asp	CTG Leu 300	GAC Asp	CCC Pro	CTG Leu	GAC Asp	912
			ACA Thr													960
			ACC Thr													1008
GGA Gly	GGC Gly	CGA Arg	GAG Glu 340	GAC Asp	CTC Leu	GTC Val	TAC Tyr	AAC Asn 345	ATC Ile	ATC Ile	TGC Cys	AAG Lys	AGC Ser 350	TGT Cys	GGC Gly	1056
			GGT Gly													1104
			CTA Leu													1152
			ACC Thr		Туг 390	Thr	Phe	Glu	Ile	Gln 395	Ala					1200
						Sano	UTITE	I E OU	וו עו	TULE	20)					

3 / 33 FIG. IC ACT GAC CAG AGC CCC TTC TCG CCT CAG TTC GCC TCT GTG AAC ATC ACC 1248 Thr Asp Gln Ser Pro Phe Ser Pro Gln Phe Ala Ser Val Asn Ile Thr 405 410 ACC AAC CAG GCA GCT CCA TCG GCA GTG TCC ATC ATG CAT CAG GTG AGC 1296 Thr Asn Gln Ala Ala Pro Ser Ala Val Ser Ile Met His Gln Val Ser 420 425 CGC ACC GTG GAC AGC ATT ACC CTG TCG TGG TCC CAG CCG GAC CAG CCC 1344 Arg Thr Val Asp Ser Ile Thr Leu Ser Trp Ser Gln Pro Asp Gln Pro 435 AAT GGC GTG ATC CTG GAC TAT GAG CTG CAG TAC TAT GAG AAG GAG CTC 1392 Asn Gly Val Ile Leu Asp Tyr Glu Leu Gln Tyr Tyr Glu Lys Glu Leu 455 AGT GAG TAC AAC GCC ACA GCC ATA AAA AGC CCC ACC AAC ACG GTC ACG 1440 Ser Glu Tyr Asn Ala Thr Ala Ile Lys Ser Pro Thr Asn Thr Val Thr 465 470 GGC CTC AAA GCC GGC GCC ATC TAT GTC TTC CAG GTG CGG GCA CGC ACT 1488 Gly Leu Lys Ala Gly Ala Ile Tyr Val Phe Gln Val Arg Ala Arg Thr 485 GTG GCA GGC TAC GGG CGC TAC AGC GGC AAG ATG TAC TTC CAG ACC ATG 1536 Val Ala Gly Tyr Gly Arg Tyr Ser Gly Lys Met Tyr Phe Gln Thr Met 505 500 ACA GAA GCC GAG TAC CAG ACA AGC ATC CAG GAG AAG TTG CCA CTC ATC 1584 Thr Glu Ala Glu Tyr Gln Thr Ser Ile Gln Glu Lys Leu Pro Leu Ile 520 ATC GGC TCC TCG GCC GCT GGC CTG GTC TTC CTC ATT GCT GTG GTT GTC 1632 Ile Gly Ser Ser Ala Ala Gly Leu Val Phe Leu Ile Ala Val Val Val 530 535 ATC GCC ATC GTG TGT AAC AGA CGG GGG TTT GAG CGT GCT GAC TCG GAG 1680 Ile Ala Ile Val Cys Asn Arg Arg Gly Phe Glu Arg Ala Asp Ser Glu 550 545 TAC ACG GAC AAG CTG CAA CAC TAC ACC AGT GGC CAC ATA ACC CCA GGC 1728 Tyr Thr Asp Lys Leu Gln His Tyr Thr Ser Gly His Ile Thr Pro Gly 565 ATG AAG ATC TAC ATC GAT CCT TTC ACC TAC GAG GAC CCC AAC GAG GCA 1776 Met Lys Ile Tyr Ile Asp Pro Phe Thr Tyr Glu Asp Pro Asn Glu Ala 580 GTG CGG GAG TTT GCC AAG GAA ATT GAC ATC TCC TGT GTC AAA ATT GAG 1824 Val Arg Glu Phe Ala Lys Glu Ile Asp Ile Ser Cys Val Lys Ile Glu 600 595

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FIG. ID CAG GTG ATC GGA GCA GGG GAG TTT GGC GAG GTC TGC AGT GGC CAC CTG 1872 Gln Val Ile Gly Ala Gly Glu Phe Gly Glu Val Cys Ser Gly His Leu 615 610 AAG CTG CCA GGC AAG AGA GAG ATC TTT GTG GCC ATC AAG ACG CTC AAG 1920 Lys Leu Pro Gly Lys Arg Glu Ile Phe Val Ala Ile Lys Thr Leu Lys 625 630 TCG GGC TAC ACG GAG AAG CAG CGC CGG GAC TTC CTG AGC GAA GCC TCC 1968 Ser Gly Tyr Thr Glu Lys Gln Arg Arg Asp Phe Leu Ser Glu Ala Ser 650 ATC ATG GGC CAG TTC GAC CAT CCC AAC GTC ATC CAC CTG GAG GGT GTC 2016 Ile Met Gly Gln Phe Asp His Pro Asn Val Ile His Leu Glu Gly Val 660 GTG ACC AAG AGC ACA CCT GTG ATG ATC ATC ACC GAG TTC ATG GAG AAT 2064 Val Thr Lys Ser Thr Pro Val Met Ile Ile Thr Glu Phe Met Glu Asn 675 680 GGC TCC CTG GAC TCC TTT CTC CGG CAA AAC GAT GGG CAG TTC ACA GTC 2112 Gly Ser Leu Asp Ser Phe Leu Arg Gln Asn Asp Gly Gln Phe Thr Val 695 ATC CAG CTG GTG GGC ATG CTT CGG GGC ATC GCA GCT GGC ATG AAG TAC 2160 Ile Gln Leu Val Gly Met Leu Arg Gly Ile Ala Ala Gly Met Lys Tyr 710 715 CTG GCA GAC ATG AAC TAT GTT CAC CGT GAC CTG GCT GCC CGC AAC ATC 2208 Leu Ala Asp Met Asn Tyr Val His Arg Asp Leu Ala Ala Arg Asn Ile 725 730 CTC GTC AAC AGC AAC CTG GTC TGC AAG GTG TCG GAC TTT GGG CTC TCA 2256 Leu Val Asn Ser Asn Leu Val Cys Lys Val Ser Asp Phe Gly Leu Ser 740 CGC TTT CTA GAG GAC GAT ACC TCA GAC CCC ACC TAC ACC AGT GCC CTG 2304 Arg Phe Leu Glu Asp Asp Thr Ser Asp Pro Thr Tyr Thr Ser Ala Leu 755 GGC GGA AAG TTC CCC ATC CGC TGG ACA GCC CCG GAA GCC ATC CAG TAC 2352 Gly Gly Lys Phe Pro Ile Arg Trp Thr Ala Pro Glu Ala Ile Gln Tvr 775 770 CGG AAG TTC ACC TCG GCC AGT GAT GTG TGG AGC TAC GGC ATT GTC ATG 2400 Arg Lys Phe Thr Ser Ala Ser Asp Val Trp Ser Tyr Gly Ile Val Met 790 795 785 TGG GAG GTG ATG TCC TAT GGG GAG CGG CCC TAC TGG GAC ATG ACC AAC 2448 Trp Glu Val Met Ser Tyr Gly Glu Arg Pro Tyr Trp Asp Met Thr Asn 805 810 SUBSTITUTE SHEET (RULE 26)

5 / 33 FIG. IE

								. 16	7. l		·					
							GAG	CAG	GAC	TAT			CCA Pro 830			2496
													TGT Cys			25,44
													AAC Asn			2592
													GCG Ala			2640
													CCC Pro			2688
													AAG Lys 910			2736
													TTT Phe			2784
													GTC Val			2832
													ATG Met			2880
						TCT Ser				TGAC	CATTO	CAC C	CTGCC	CTCGO	SC .	2930
TCAC	CTCT	rrc (CTCCA	AAGC	cc co	GCCC	CCTCT	r GC								2962

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FIG. 2A

						•	r	-10). C	ZΑ						
															CTC Leu	48
		TGC Cys														96
		AAC Asn 35														144
		TGG Trp														192
		GAA Glu														240
		CAG Gln														288
		GCT Ala														336
		AGC Ser 115														384
		TAC Tyr														432
		TAC Tyr														480
GAA Glu	CTT Leu	GAT Asp	CTT Leu	GGT Gly 165	GAC Asp	CGT Arg	GTT Val	ATG Met	AAA Lys 170	CTG Leu	AAT Asn	ACA Thr	GAG Glu	GTC Val 175	AGA Arg	528
		GGA Gly														576
		GCT Ala 195		Ile	Ala	Leu	Val 200	Ser	Val	Arg						624
				C	רסמוני	CITIITI	ב כאב	FT (R	HIF?	/61						

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FIG. 2B

						- 1	- 1(j. 7	길B				
					CAC His 215	TTG	GCT	GTC	TTC				672
(TTG Leu								720
					CCT Pro								768
					GGG Gly					\			816
					CAA Gln								864
					TGC Cys 295								912
1					TCT Ser								960
					ACA Thr								1008
					AAT Asn							,	1056
					ACT Thr								1104
					AAC Asn 375								1152
(CTT Leu								1200
			Val 405	Asp	CTA Leu	Leu	Ala	His 410	Thr				1248
				יו אוווי:	C) T1 1T1	CUL	r T / 17	1 11 F 7	(C')				

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8 / 33 FIG. 2C ATT GAG GCA GTG AAT GGA GTG TCC GAC TTG AGC CCA GGA GCC CGG CAG 1296 Ile Glu Ala Val Asn Gly Val Ser Asp Leu Ser Pro Gly Ala Arg Gln 420 425 TAT GTG TCT GTA AAT GTA ACC ACA AAT CAA GCA GCT CCA TCT CCA GTC 1344 Tyr Val Ser Val Asn Val Thr Thr Asn Gln Ala Ala Pro Ser Pro Val 435 ACC AAT GTG AAA AAA GGG AAA ATT GCA AAA AAC AGC ATC TCT TTG TCT 1392 Thr Asn Val Lys Lys Gly Lys Ile Ala Lys Asn Ser Ile Ser Leu Ser 455 460 TGG CAA GAA CCA GAT CGT CCC AAT GGA ATC ATC CTA GAG TAT GAA ATC 1440 Trp Gln Glu Pro Asp Arg Pro Asn Gly Ile Ile Leu Glu Tyr Glu Ile 465 470 AAG CAT TTT GAA AAG GAC CAA GAG ACC AGC TAC ACG ATT ATC AAA TCT 1488 Lys His Phe Glu Lys Asp Gln Glu Thr Ser Tyr Thr Ile Ile Lys Ser 485 490 AAA GAG ACA ACT ATT ACT GCA GAG GGC TTG AAA CCA GCT TCA GTT TAT 1536 Lys Glu Thr Thr Ile Thr Ala Glu Gly Leu Lys Pro Ala Ser Val Tyr 500 505 GTC TTC CAA ATT CGA GCA CGT ACA GCA GCA GGC TAT GGT GTC TTC AGT 1584 Val Phe Gln Ile Arg Ala Arg Thr Ala Ala Gly Tyr Gly Val Phe Ser 515 520 CGA AGA TTT GAG TTT GAA ACC ACC CCA GTG TTT GCA GCA TCC AGC GAT 1632 Arg Arg Phe Glu Phe Glu Thr Thr Pro Val Phe Ala Ala Ser Ser Asp 530 535 540 CAA AGC CAG ATT CCT GTA ATT GCT GTG TCT GTG ACA GTA GGA GTC ATT 1680 Gln Ser Gln Ile Pro Val Ile Ala Val Ser Val Thr Val Gly Val Ile 545 550 TTG TTG GCA GTG GTT ATC GGC GTC CTC CTC AGT GGA AGG CGG TGT GGC 1728 Leu Leu Ala Val Val Ile Gly Val Leu Leu Ser Gly Arg Arg Cys Gly 565 570 TAC AGC AAA GCA AAA CAA GAT CCA GAA GAG GAA AAG ATG CAT TTT CAT 1776 Tyr Ser Lys Ala Lys Gln Asp Pro Glu Glu Glu Lys Met His Phe His 580 585 AAT GGG CAC ATT AAA CTG CCA GGA GTA AGA ACT TAC ATT GAT CCA CAT 1824 Asn Gly His Ile Lys Leu Pro Gly Val Arg Thr Tyr Ile Asp Pro His 600 ACC TAT GAG GAT CCC AAT CAA GCT GTC CAC GAA TTT GCC AAG GAG ATA 1872 Thr Tyr Glu Asp Pro Asn Gln Ala Val His Glu Phe Ala Lys Glu Ile 610 615 620 SUBSTITUTE SHEET (RULE 26)

9 / 33 FIG. 2D GAA GCA TCA TGT ATC ACC ATT GAG AGA GTT ATT GGA GCA GGT GAA TTT 1920 Glu Ala Ser Cys Ile Thr Ile Glu Arg Val Ile Gly Ala Gly Glu Phe 630 GGT GAA GTT TGT AGT GGA CGT TTG AAA CTA CCA GGA AAA AGA GAA TTA 1968 Gly Glu Val Cys Ser Gly Arg Leu Lys Leu Pro Gly Lys Arg Glu Leu 645 · 650 CCT GTG GCT ATC AAA ACC CTT AAA GTA GGC TAT ACT GAA AAG CAA CGC 2016 Pro Val Ala Ile Lys Thr Leu Lys Val Gly Tyr Thr Glu Lys Gln Arg AGA GAT TTC CTA GGT GAA GCA AGT ATC ATG GGA CAG TTT GAT CAT CCT 2064 Arg Asp Phe Leu Gly Glu Ala Ser Ile Met Gly Gln Phe Asp His Pro 680 AAC ATC ATC CAT TTA GAA GGT GTG GTG ACC AAA AGT AAA CCA GTG ATG 2112 Asn Ile Ile His Leu Glu Gly Val Val Thr Lys Ser Lys Pro Val Met 690 695 ATC GTG ACA GAG TAT ATG GAG AAT GGC TCT TTA GAT ACA TTT TTG AAG 2160 Ile Val Thr Glu Tyr Met Glu Asn Gly Ser Leu Asp Thr Phe Leu Lys 705 710 720 AAA AAC GAT GGG CAG TTC ACT GTG ATT CAG CTT GTT GGC ATG CTG AGA 2208 Lys Asn Asp Gly Gln Phe Thr Val Ile Gln Leu Val Gly Met Leu Arg 725 GGT ATC TCT GCA GGA ATG AAG TAC CTT TCT GAC ATG GGC TAT GTG CAT 2256 Gly Ile Ser Ala Gly Met Lys Tyr Leu Ser Asp Met Gly Tyr Val His 740 745 AGA GAT CTT GCT GCC AGA AAC ATC TTA ATC AAC AGT AAC CTT GTG TGC 2304 Arg Asp Leu Ala Ala Arg Asn Ile Leu Ile Asn Ser Asn Leu Val Cys 760 AAA GTG TCT GAC TTT GGA CTT TCC CGG GTA CTG GAA GAT GAT.CCC GAG 2352 Lys Val Ser Asp Phe Gly Leu Ser Arg Val Leu Glu Asp Asp Pro Glu 770 775 GCA GCC TAC ACC ACA AGG GGA GGA AAA ATT CCA ATC AGA TGG ACT GCC 2400 Ala Ala Tyr Thr Thr Arg Gly Gly Lys Ile Pro Ile Arg Trp Thr Ala 785 800 CCA GAA GCA ATA GCT TTC CGA AAG TTT ACT TCT GCC AGT GAT GTC TGG 2448 Pro Glu Ala Ile Ala Phe Arg Lys Phe Thr Ser Ala Ser Asp Val Trp 805 810 815 AGT TAT GGA ATA GTA ATG TGG GAA GTT GTG TCT TAT GGA GAG AGA CCC 2496 Ser Tyr Gly Ile Val Met Trp Glu Val Val Ser Tyr Gly Glu Arg Pro 820 825

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10/33 FIG. 2E																
TAC Tyr	TGG Trp	GAG Glu 835	ATG Met	ACC Thr	AAT Asn	CAA Gln	GAT	GTG	ATT	AAA	GCG	GTA Val 845	GÀG Glu	GAA Glu	GGC Gly	254
TAT Tyr	CGT Arg 850	CTG Leu	CCA Pro	AGC Ser	CCC Pro	ATG Met 855	GAT Asp	TGT Cys	CCT Pro	GCT Ala	GCT Ala 860	CTC Leu	TAT Tyr	CAG Gln	TTA Leu	2592
ATG Met 865	CTG Leu	GAT Asp	TGC Cys	TGG Trp	CAG Gln 870	AAA Lys	GAG Glu	CGA Arg	AAT Asn	AGC Ser 875	AGG Arg	CCC Pro	AAG Lys	TTT Phe	GAT Asp 880	2640
			AAC Asn													2688
AAG Lys	ACG Thr	CTG Leu	GTT Val 900	AAT Asn	GCA Ala	TCC Ser	TGC Cys	AGA Arg 905	GTA Val	TCT Ser	AAT Asn	TTA Leu	TTG Leu 910	GCA Ala	GAA Glu	2736
			CTA Leu													2784
		Ile	AAG Lys													2832
			ATG Met													2880
			GTG Val													2928
2983			ATG Met 980				Leu								TAACTTCA	rg
TAAA	TGTC	GC I	TCTT	'CAAG	T GA	ATGA	TTCT	' GCA	CTTI	GTA	AACA	GCAC	TG A	GATT.	TATTT	3043
TAAC	AAAA	AA A	GGGG	GAAA	A GG	GAAA	ACAG	TGA	TTTC	TAA	ACCT	TAGA	AA A	CATT	TGCCT	3103
ርልGCCACAGA												ChCth	3162			

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							F	=1()	3 A							
AAGCGGCAGG AGCAGCGTTG GCACCGGCGA ACC ATG GCT GGG ATT TTC TAT TTC Met Ala Gly Ile Phe Tyr Phe 1 5 GCC CTA TTT TCG TGT CTC TTC GGG ATT TGC GAC GCT GTC ACA GGT TCC															54		
														GGT Gly			102
AGG Arg	GTA Val 25	TAC Tyr	CCC Pro	GCG Ala	AAT Asn	GAA Glu 30	GTT Val	ACC Thr	TTA Leu	TTG Leu	GAT Asp 35	TCC Ser	AGA Arg	TCT Ser	GTT Val		150
														TGG Trp			198
														TAC Tyr 70			246
														ACT Thr			294
														AAA Lys			342
														TGC Cys			390
														GAG Glu			438
														GCT Ala 150			486
														CTG Leu			534
														TAC Tyr			582

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12/33 FIG. 3B GCT TTT CAG GAT GTG GGG GCC TGC ATC GCC CTG GTA TCA GTC CGT GTG 630 Ala Phe Gln Asp Val Gly Ala Cys Ile Ala Leu Val Ser Val Arg Val 185 190 TTC TAT AAA AAG TGT CCA CTC ACA GTC CGC AAT CTG GCC CAG TTT CCT 678 Phe Tyr Lys Lys Cys Pro Leu Thr Val Arg Asn Leu Ala Gln Phe Pro 200 205 GAC ACC ATC ACA GGG GCT GAT ACG TCT TCC CTG GTG GAA GTT CGA GGC 726 Asp Thr Ile Thr Gly Ala Asp Thr Ser Ser Leu Val Glu Val Arg Gly 220 225 TCC TGT GTC AAC AAC TCA GAA GAG AAA GAT GTG CCA AAA ATG TAC TGT 774 Ser Cys Val Asn Asn Ser Glu Glu Lys Asp Val Pro Lys Met Tyr Cys GGG GCA GAT GGT GAA TGG CTG GTA CCC ATT GGC AAC TGC CTA TGC AAC 822 Gly Ala Asp Gly Glu Trp Leu Val Pro Ile Gly Asn Cys Leu Cys Asn 255 GCT GGG CAT GAG GAG CGG AGC GGA GAA TGC CAA GCT TGC AAA ATT GGA 870 Ala Gly His Glu Glu Arg Ser Gly Glu Cys Gln Ala Cys Lys Ile Gly 265 270 TAT TAC AAG GCT CTC TCC ACG GAT GCC ACC TGT GCC AAG TGC CCA CCC 918 Tyr Tyr Lys Ala Leu Ser Thr Asp Ala Thr Cys Ala Lys Cys Pro Pro 280 295 CAC AGC TAC TCT GTC TGG GAA GGA GCC ACC TCG TGC ACC TGT GAC CGA 966 His Ser Tyr Ser Val Trp Glu Gly Ala Thr Ser Cys Thr Cys Asp Arg 300 305 GGC TTT TTC AGA GCT GAC AAC GAT GCT GCC TCT ATG CCC TGC ACC CGT 1014 Gly Phe Phe Arg Ala Asp Asn Asp Ala Ala Ser Met Pro Cys Thr Arg 315 320 CCA CCA TCT GCT CCC CTG AAC TTG ATT TCA AAT GTC AAC GAG ACA TCT 1062 Pro Pro Ser Ala Pro Leu Asn Leu Ile Ser Asn Val Asn Glu Thr Ser 335 GTG AAC TTG GAA TGG AGT AGC CCT CAG AAT ACA GGT GGC CGC CAG GAC 1110 Val Asn Leu Glu Trp Ser Ser Pro Gln Asn Thr Gly Gly Arg Gln Asp 350 345 ATT TCC TAT AAT GTG GTA TGC AAG AAA TGT GGA GCT GGT GAC CCC AGC 1158 Ile Ser Tyr Asn Val Val Cys Lys Cys Gly Ala Gly Asp Pro Ser 360 365 375 AAG TGC CGA CCC TGT GGA AGT GGG GTC CAC TAC ACC CCA CAG CAG AAT 1206 Lys Cys Arg Pro Cys Gly Ser Gly Val His Tyr Thr Pro Gln Gln Asn

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13 / 33 FIG. 3C

							ŀ	- [(э. J	3C							
GGC Gly	TTG Leu	AAG Lys	ACC Thr 395	Thr	AAA Lys	GTC Val	TCC	ATC	ACT Thr	GAC	CTC Leu	CTA Leu	GCT Ala 405	His	ACC Thr		1254
AAT Asn	TAC Tyr	ACC Thr 410	TTT Phe	GAA Glu	ATC Ile	TGG Trp	GCT Ala 415	GTG Val	AAT Asn	GGA Gly	GTG Val	TCC Ser 420	AAA Lys	TAT	AAC Asn		1302
CCT Pro	AAC Asn 425	CCA Pro	GAC Asp	CAA Gln	TCA Ser	GTT Val 430	TCT Ser	GTC Val	ACT Thr	GTG Val	ACC Thr 435	ACC Thr	AAC Asn	CAA Gln	GCA Ala	. ···	1350
GCA Ala 440	CCA Pro	TCA Ser	TCC Ser	ATT Ile	GCT Ala 445	TTG Leu	GTC Val	CAG Gln	GCT Ala	AAA Lys 450	GAA Glu	GTC Val	ACA Thr	AGA Arg	TAC Tyr 455		1398
AGT Ser	GTG Val	GCA Ala	CTG Leu	GCT Ala 460	TGG Trp	CTG Leu	GAA Glu	CCA Pro	GAT Asp 465	CGG Arg	CCC Pro	AAT Asn	GGG Gly	GTA Val 470	ATC Ile		1446
CTG Leu	GAA Glu	TAT Tyr	GAA Glu 475	GTC Val	AAG Lys	TAT Tyr	TAT Tyr	GAG Glu 480	AAG Lys	GAT Asp	CAG Gln	AAT Asn	GAG Glu 485	CGA Arg	AGC Ser		1494
TAT Tyr	CGT Arg	ATA Ile 490	GTT Val	CGG Arg	ACA Thr	GCT Ala	GCC Ala 495	AGG Arg	AAC Asn	ACA Thr	GAT Asp	ATC Ile 500	AAA Lys	GGC Gly	CTG Leu		1542
AAC Asn	CCT Pro 505	CTC Leu	ACT Thr	TCC Ser	TAT Tyr	GTT Val 510	TTC Phe	CAC His	GTG Val	CGA Arg	GCC Ala 515	AGG Arg	ACA Thr	GCA Ala	GCT Ala		1590
												ACC Thr					1638
												GTC Val					1686
												ATT Ile					1734
												AAA Lys 580					1782
				His	Leu	Asn 590	Gln	Gly				TAT Tyr					1830
				CH	DOTIN	1177	11 IF F-1										

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14/33 FIG. 3D TTT ACG TAC GAA GAT CCC AAC CAA GCA GTG CGA GAG TTT GCC AAA GAA 1878 Phe Thr Tyr Glu Asp Pro Asn Gln Ala Val Arg Glu Phe Ala Lys Glu 605 ATT GAC GCA TCC TGC ATT AAG ATT GAA AAA GTT ATA GGA GTT GGT GAA 1926 Ile Asp Ala Ser Cys Ile Lys Ile Glu Lys Val Ile Gly Val Gly Glu 620 -TTT GGT GAG GTA TGC AGT GGG CGT CTC AAA GTG CCT GGC AAG AGA GAG 1974 Phe Gly Glu Val Cys Ser Gly Arg Leu Lys Val Pro Gly Lys Arg Glu 640 ATC TGT GTG GCT ATC AAG ACT CTG AAA GCT GGT TAT ACA GAC AAA CAG 2022 Ile Cys Val Ala Ile Lys Thr Leu Lys Ala Gly Tyr Thr Asp Lys Gln 655 AGG AGA GAC TTC CTG AGT GAG GCC AGC ATC ATG GGA CAG TTT GAC CAT 2070 Arg Arg Asp Phe Leu Ser Glu Ala Ser Ile Met Gly Gln Phe Asp His 670 675 CCG AAC ATC ATT CAC TTG GAA GGC GTG GTC ACT AAA TGT AAA CCA GTA 2118 Pro Asn Ile Ile His Leu Glu Gly Val Val Thr Lys Cys Lys Pro Val 680 685 ATG ATC ATA ACA GAG TAC ATG GAG AAT GGC TCC TTG GAT GCA TTC CTC 2166 Met Ile Ile Thr Glu Tyr Met Glu Asn Gly Ser Leu Asp Ala Phe Leu 700 AGG AAA AAT GAT GGC AGA TTT ACA GTC ATT CAG CTG GTG GGC ATG CTT 2214 Arg Lys Asn Asp Gly Arg Phe Thr Val Ile Gln Leu Val Gly Met Leu 715 720 CGT GGC ATT GGG TCT GGG ATG AAG TAT TTA TCT GAT ATG AGC TAT GTG 2262 Arg Gly Ile Gly Ser Gly Met Lys Tyr Leu Ser Asp Met Ser Tyr Val CAT CGT GAT CTG GCC GCA CGG AAC ATC CTG GTG AAC AGC AAC TTG GTC 2310 His Arg Asp Leu Ala Ala Arg Asn Ile Leu Val Asn Ser Asn Leu Val 750 745 TGC AAA GTG TCT GAT TTT GGC ATG TCC CGA GTG CTT GAG GAT GAT CCG 2358 Cys Lys Val Ser Asp Phe Gly Met Ser Arg Val Leu Glu Asp Asp Pro 760 765 770 GAA GCA GCT TAC ACC ACC AGG GGT GGC AAG ATT CCT ATC CGG TGG ACT 2406 Glu Ala Ala Tyr Thr Thr Arg Gly Gly Lys Ile Pro Ile Arg Trp Thr 789

800 SUBSTITUTE SHEET (RULE 26)

GCG CCA GAA GCA ATT GCC TAT CGT AAA TTC ACA TCA GCA AGT GAT GTA

Ala Pro Glu Ala Ile Ala Tyr Arg Lys Phe Thr Ser Ala Ser Asp Val

15 / 33 FIG. 3E																
							TGG	GAA	GTG	ATG				GAG Glu		2502
														GAG Glu		2550
														CAC His		2598
														AAA Lys 870		2646
														AAC Asn		2694
														TTG Leu		2742
														GAT Asp		2790
														GCT Ala		2838
				Leu		Ala	Val	Val	His	Val		Gln		GAC Asp 950	Leu	2886
		Ile					Ile							TTG Leu		2934
														AGA Arg		2982
	CCC Pro 985		TGAG	CCAC	STA (TGAF	ATAA?	C TC	CAAA <i>P</i>	CTCI	TGA	LTAA	'AGT			3031
TTAC	CTC	ATC C	CATGC	CACTI	T A	ATTG <i>I</i>	\AGA#	CTO	CACT	TTT	TTTA	CTTC	GT C	TTCG	CCCTC	3091
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16/33 FIG 4Δ

				•						ナム							
CGGT	'GCGZ	AGC (GAAC	AGGA	GT G	GGGG	GGAA	A TT	AAAA.	AAAG	CTA	AACG'	TGG .	AGCA	GCCGAT	ľ	60
CGGG	GAC	CGA (GAAG(GGGA	AT C	GATG	CAAG	G AG	CACA	СТАА	AAC	AAAA	GCT .	ACTT	CGGAA	2	120
AAAC	AGC	ATT ?	IAAA1	AATC	CA CO	GACT	CAAG	A TAI	ACTG	AAAC	CTA	AAAT	AAA /	ACCT(GCTCAT	ŗ	180
GCAC										er Ti				TA TO eu Cy			227
															GCG Ala 30		275
														TTG Leu 45			323
														TTG Leu			371
														ATG Met			419
														GGC Gly			467
														TGT Cys			515
														TTG Leu 125			563
														AAC Asn			611
														CAA Gln			659
Asp :														GAG Glu			707

17/33 FIG. 4B

				- F	- 16	j. Z	18						
											GGG Gly 190		755
									AAG Lys				803
									GTG Val				851
									GTC Val 235				899
									AGT Ser				947
									GCA Ala				995
									TTC Phe			1	043
									CAC His			1	091
									GGG Gly 315			1	139
									CCT Pro			1	187
									GTA Val			1	235
									GTG Val			. 1	283
			Ser	Trp	Glu 375	Gln	Gly	Glu	TGT Cys			1	331
			20B	UIIIC	TE SH	ובבו (HULL	20)					

18/33 FIG. 40

FIG. 4C														
AGT Ser						CCC	CAG	CAG	ACT					1379
GTC Val 400	Thr													1427
GAA Glu														1475
GCT Ala														1523
GGA Gly														 1571
CAG Gln														 1619
TAT Tyr 480														1667
AAG Lys														1715
GTT Val														1763
CCC Pro														1811
GAA Glu														1859
GTG Val 560														1907
TTC Phe				Arg 580		His	Cys	Gly	Tyr 585	Ser				1955

^{19/33} FIG 4D

¹⁹⁷³³ FIG. 4D																
				GAG	CTT	TAC Tyr	TTT	CAT	TTT	AAA						2003
						ACC Thr										2051
						GAT Asp						,				2099
						GGT Gly 645										2147
						GCA Ala										2195
						AGA Arg										2243
						AAT Asn										2291
						ATA Ile										2339
						AAA Lys 725										2387
TTA Leu 735	GTA Val	GGA Gly	ATG Met	CTG Leu	AGA Arg 740	GGA Gly	ATT Ile	GCT Ala	GCT Ala	GGA Gly 745	ATG Met	AGA Arg	TAT Tyr	TTG Leu	GCT Ala 750	2435
GAT Asp	ATG Met	GGA Gly	TAT Tyr	GTT Val 755	CAC His	AGG Arg	GAC Asp	CTT Leu	GCA Ala 760	GCT Ala	CGC Arg	AAT Asn	ATT Ile	CTT Leu 765	GTC Val	2483
AAC Asn	AGC Ser	AAT Asn	CTC Leu 770	GTT Val	TGT. Cys	AAA Lys	GTG Val	TCA Ser 775	GAT Asp	TTT Phe	GGC Gly	CTG Leu	TCC Ser 780	CGA Arg	GTT Val	2531
ATA Ile	GAG Glu	GAT Asp 785	GAT Asp	CCA Pro	Glu	GCT Ala IBSTI	Val 790	Tyr	Thr	Thr	Thr	GGT Gly 795	GGA Gly	AAA Lys	ATT Ile	2579
					ĢU	ווטטו		J. 166			•					

^{20/33} FIG. 4E CCA GTA AGG TGG ACA GCA CCC GAA GCC ATC CAG TAC CGG AAA TTC ACA 2627 Pro Val Arg Trp Thr Ala Pro Glu Ala Ile Gln Tyr Arg Lys Phe Thr 805 810 800 TCA GCC AGT GAT GTA TGG AGC TAT GGA ATA GTC ATG TGG GAA GTT ATG 2675 Ser Ala Ser Asp Val Trp Ser Tyr Gly Ile Val Met Trp Glu Val Met 815 825 TCT TAT GGA GAA AGA CCT TAT TGG GAC ATG TCA AAT CAA GAT GTT ATA 2723 Ser Tyr Gly Glu Arg Pro Tyr Trp Asp Met Ser Asn Gln Asp Val Ile 840 845 835 2771 AAA GCA ATA GAA GAA GGT TAT CGT TTA CCA GCA CCC ATG GAC TGC CCA Lys Ala Ile Glu Glu Gly Tyr Arg Leu Pro Ala Pro Met Asp Cys Pro 850 855 GCT GGC CTT CAC CAG CTA ATG TTG GAT TGT TGG CAA AAG GAG CGT GCT 2819 Ala Gly Leu His Gln Leu Met Leu Asp Cys Trp Gln Lys Glu Arg Ala 870 865 2867 GAA AGG CCA AAA TTT GAA CAG ATA GTT GGA ATT CTA GAC AAA ATG ATT Glu Arg Pro Lys Phe Glu Gln Ile Val Gly Ile Leu Asp Lys Met Ile 885 880 CGA AAC CCA AAT AGT CTG AAA ACT CCC CTG GGA ACT TGT AGT AGG CCA 2915 Arg Asn Pro Asn Ser Leu Lys Thr Pro Leu Gly Thr Cys Ser Arg Pro 900 905 ATA AGC CCT CTT CTG GAT CAA AAC ACT CCT GAT TTC ACT ACC TTT TGT 2963 Ile Ser Pro Leu Leu Asp Gln Asn Thr Pro Asp Phe Thr Thr Phe Cys 920 915 TCA GTT GGA GAA TGG CTA CAA GCT ATT AAG ATG GAA AGA TAT AAA GAT 3011 Ser Val Gly Glu Trp Leu Gln Ala Ile Lys Met Glu Arg Tyr Lys Asp 930 935 AAT TTC ACG GCA GCT GGC TAC AAT TCC CTT GAA TCA GTA GCC AGG ATG 3059 Asn Phe Thr Ala Ala Gly Tyr Asn Ser Leu Glu Ser Val Ala Arg Met 955 950 945 ACT ATT GAG GAT GTG ATG AGT TTA GGG ATC ACA CTG GTT GGT CAT CAA 3107 Thr Ile Glu Asp Val Met Ser Leu Gly Ile Thr Leu Val Gly His Gln 965 960 AAG AAA ATC ATG AGC AGC ATT CAG ACT ATG AGA GCA CAA ATG CTA CAT 3155 Lys Lys Ile Met Ser Ser Ile Gln Thr Met Arg Ala Gln Met Leu His 985 980 TTA CAT GGA ACT GGC ATT CAA GTG TGATATGCAT TTCTCCCTTT TAAGGGAGAT 3209 Leu His Gly Thr Gly Ile Gln Val

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FIG. 4F

TACAGACT	GC AA	GAGAACAG	TACTGGCCTT	CAGTATATGC	ATAGAATGCT	GCTAGAAGAC	3269
AAGTGATG'	TC CT	GGGTCCTT	CCAACAGTGA	AGAGAAGATT	TAAGAAGCAC	CTATAGACTT	3329
GAACTCCT	AA GT	GCCACCAG	ААТАТАТАА	AAGGGAATTT	AGGATCCACC	ATCGGTGGCC	3389
AGGAAAAT	AG CA	GTGACAAT	AAACAAAGTA	CTACCTGAAA	AACATCCAAA	CACCTTGAGC	3449
TCTCTAAC	CT CC	TTTTTGTC	TTATAGACTT	TTTAAAATGT	ACATAAAGAA	TTTAAGAAAG	3509
AATATATT'	TG TC	AAATAAAA	TCATGATCTT	ATTGTTAAAA	TTAATGAAAT	ATTTTCCTTA	3569
AATATGTG2	AT TT	CAGACTAT	TCCTTTTTAA	AATCATTTGT	GTTTATTCTT	CATAAGGACT	3629
TTGTTTTA(GA AA	GCTGTTTA	TAGCTTTGGA	CCTTTTTAGT	GTTAAATCTG	TAACATTACT	3689
ACACTGGG'	ra cc'	TTTGAAAG	AATCTCAAAT	TTCAAAAGAA	ATAGCATGAT	TGAAGATACA	3749
TCTCTGTT	AG AA	CATTGGTA	TCCTTTTTGT	GCCATTTTAT	TCTGTTTAAT	CAGTGCTGTT	3809
TTGATATT(GT TT	GCTAATTG	GCAGGTAGTC	AAGAAAATGC	AAGTTGCCAA	GAGCTCTGAT	3869
ATTTTTTA	AA AA	GAATTTTT	TTGTAAAGAT	CAGACAACAC	ACTATCTTTT	CAATGAAAAA	3929
AGCAATAAT	rg at	CCATACAT	ACTATAAGGC	ACTTTTAACA	GATTGTTTAT	AGAGTGATTT	3989
TACTAGAAA	AG AA'	PTTAATAA	ACTCGAAGTT	TAGGTTTATG	AGTATATAAA	CAAATGAGGC	4049
ACTTCATCI	rg aa	GAATGTTG	GTGAAGGCAA	GTCTCTGAAA	GCAGAACTAT	CCAGTGTTAT	4109
CTAAAAATT	TA AT	CTGAGCAC	ATCAAGATTT	TTTCATTCTC	GTGACATTAG	GAAATTTAGG	4169
ATAAATAGT	TT GAG	CATATATT	TTATATCCTC	TTCTGTTGAA	TGCAGTCCAA	ACATGAAAGG	4229
AAATAATTO	ST TT	TATATAT	AACTCTGAAG	CATGATAAAG	GGGCAGTTCA	CAATTTTCAC	4289
CATTTAAAC	CA CA	AATTTGCT	GCACAGAATA	TCACCATTGC	AGTTCAAAAC	AAAACAAAAC	4349
AAAAAGTCI	TT TT	GTTTGTGA	ACACTGATGC	AAGAAACTTG	TTAAATGAAA	GGACTCTTTA	4409
CCCTAGAAG	G AA	GAGGTGAA	GGATCTGGCT	TGTTTTTAAA	GCTTTATTTA	ТТАААССАТА	4469
TTATTTGAT	T AC	TGTGTTAG	AATTTCATAA	GCAATAATTA	AATGTGTCTT	TATGGAATTC	4529

FIG. 5A

11 2 8 2 2 1 1 2	🕏 EPH 🥏 RGEEASRVHVELQFTVRDCKSFPGGAGPLGCKETFNLLYMESDQDVGIQLRRPLFQKVTTVAADQSFTIRDLASGSVKLNVERCSLGRLTRRGLYL	ECK RG.EAERNNFELNFTVRDCNSFPGGASSCKETFNLYYAESDLDYGTNFQKRLFTKIDTIAPDEITVSSDFEARHVKLNVEERSVGPLTRKGFYL	HEK4 RN.SAQKIYVELKFTLRDCNSIPLVLGTCKETFNLYYMESDDDHGVKFREHQFTKIDTIAADESFTQMDLGDRILKLNTEIREVGPVNKKGFYL	HEKS RR.GAHRIHVEMKFSVRDCSSIPSVPGSCKETFNLYYYEADFDSATKTFPNWMENPWVKVDTIAADESFSQVDLGGRVMKINTEVRSFGPVSRSGFYL	HEK7 NE.GASRIFIELKFTLRDCNSLPGGLGTCKETFNMYYFESDDQNGRNIKENQYIKIDTIAADESFTELDLGDRVMKLNTEVRDVGPLSKKGFYL	HEK8 RE.GAQRVYIEIKFTLRDCNSLPGVMGTCKETFNLYYYESDNDKERFIRENQFVKIDTIAADESFTQVDIGDRIMKLNTEIRDVGPLSKKGFYL	HEK2 RR.DVQRVYVELKFTVRDCNSIPNIPGSCKETFNLFYYEADSDVASASSPFWMENPYVKVDTIAPDESFSRLDAGRVNTKVRSFGPLSKAGFYL	HEK11 KG.NAORIFVELKFTLRDCNSLPGVLGT. CKETFNLYYYETDYD TGRNTRENLYVKTDTTAADRSFTOGDLGFRKMKLNTFVPFTGDLGORGEVT.
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FIG. 5B

..VTDEPPKMHCSAEGEWLVPIGKCMCKAGYEEK.NGT.CQVCR SGTFKANQGDEACTHCPINSRTTSEGATNCVCRNGYYRADLDPLDMPCTTIPSAPQAVISSVNETSLMLEWTPPRDSGGREDLVYNIICKSCGSGR....G AFgdvGaC.aLvsVrv.ykkCpstv.nlA.FpdT.tgadsssLvevrG.Cvnna....e...pp.m.CsadGEWlVPiGkC.CkaGyee...gtaCqaCp AFHNPGACVALVSVRVFYQRCPETLNGLAQFPDTLPG. PA. GLVEVAGTCLPHARASPRPSGAPRMHCSPDGEWLVPVGRCHCEPGYEEGGSGEACVACP AFQDIGACVALLSVRVYYKKCPELLQGLAHFPETIAGSDAPSLATVAGTCVDHA.VVPPGGEEPRMHCAVDGEWLVPIGQCLCQAGYEKVED..ACQACS ..KEEDPPRMYCSTEGEWLVPIGKCSCNAGYEER,.GFMCQACR .. EEVDVPIKLYCNGDGEWLVPIGRCMCKAGFEAVENGTVCRGCP .. EEKDVPKMYCGADGEWLVPIGNCLCNAGHEER.. SGECQACK AFQDQGACMSLISVRAFYKKCASTTAGFALFPETLTGAEPTSLVIAPGTCIPNA...VEVSVPLKLYCNGDGEWMVPVGACTCATGHEPAAKESQCRPCP AFQDVGACIALVSVKVYYKKCWSIIENLAIFPDTVTGSEFSSLVEVRGTCVSSA..EEEAENAPRMHCSAEGEWLVPIGKCICKAGYQQK..GDTCEPCG pGfyka..gd.pClkCPphs.ttsegatsCtCengy.RadsdppsmaCTrpPSaPrnlisnvnetsv.LeWspPadtGgR.Dv.yn.iCkkCg.ga...g SGSYRMDMDTPHCLTCPQQSTAESEGATICTCESGHYRAPGEGPQVACTGPPSAPRNLSFSASGTQLSLRWEPPADTGGRQDVRYSVRCSQCQGTAQDGG PGFFKFEASESPCLECPEHTLPSPEGATSCECEEGFFRAPQDPASMPCTRPPSAPHYLTAVGMGAKVELRWTPPQDSGGREDIVYSVTCEQCWPES...G PGFYKALDGNMKCAKCPPHSSTQEDGSMNCRCENNYFRADKDPPSMACTRPPSSPRNVISNINETSVILDWSWPLDTGGRKDVTFNIICKKCGWNI...K PGFFKASPHIQSCGKCPPHSYTHEEASTSCVCEKDYFRRESDPPTMACTRPPSAPRNAISNVNETSVFLEWIPPADTGGRKDVSYYIACKKCNSHA...G IGYYKALSTDATCAKCPPHSYSVWEGATSCTCDRGFFRADNDAASMPCTRPPSAPLNLISNVNETSVNLEWSSPQNTGGRQDISYNVVCKKCGAGD..PS PGSYKAKQGEGPCLPCPPNSRTTSPAASICTCHNNFYRADSDSADSACTTVPSPPRGVISNVNETSLILEWSEPRDLGVRDDLLYNVICKKC.HGAGGAS RGFYKSSSQDLQCSRCPTHSFSDKEGSSRCECEDGYYRAPSDPPYVACTRPPSAPQNLIFNINQTTVSLEWSPPADNGGRNDVTYRIİCKRCSWEQ...G AFQDYGGCMSLIAVRVFYRKCPRIIQNGAIFQETLSGAESTSLVAARGSCIANA. AFQDVGACIALVSVRVFYKKCPLTVRNLAQFPDTITGADTSSLVEVRGSCVNNS. AFQDVGACIALVSVRVYYKKCPSVVRHLAVFPDTITGADSSQLLEVSGSCVNHS. AFQDVGACVALVSVRVYFKKCPFTVKNLAMFPDTVP.MDSQSLVEVRGSCVNNS. HEK11 HEK8 CONS HEK5 HEK2 HEK4 HEK7 CONS HEK4 HEK5 TEK8 HEK7 HEK2 EPH ECK EPH ECK SUBSTITUTE SHEET (RULE 26)

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LEYEVKYYEKDQNERSYRIVRTAARNTDIKGLNPLTSYVFHVRARTAAGYGDFSEPLEVTTNTVPSRIIGDGANSTVLLVSVSGSVVLVVILIAAFVIS [TEYEIKYYEKDQRERTYSTVKTKSTSASINNLKPGTVYVPQIRAFTAAGYGNYSPRLDVATLEEATGKMFEATAVSSEQNPVIIIAVVXVAGTIILVFM NLTYE....LHVLNQDEERYQMVLEPRVLLTELQPDTTYIVRVRMLTPLGPGPFSPDHEFRTSPPVSRGLTGGEIVAVIFGLLLGAALLLGILVFRSRRA ILDYEVKYYEKQEQETSYTILRARGINVTISSLKPDTIYVLQIRARTAAGYGTNSRKFEFETSPDSFSISGESSQVVMIAISAAVAIILLTVVIYVLIGR :LDYELQYYEKELSEYNATAIKSPTNTVTVQGLKAGAIYVFQVRARTVAGYGRYSGKMYFQTMTEAEYQTSIQEKLPLIIGSSAAGLVFLIAVVVIAIVC LEYEIKHFEKDQETSYTII.KSKETTITAEGLKPASVYVFQIRARTAAGYGVFSRRFEFETTPVFAASSDQSQIPVIAVSVTVGVILLAVVIGVLLSGR LDYEMKYFEK..SEGIASTVTSQMNSVQLDGLRPDARYVVQVRARTVAGYGQYSRPAEFETTSERGSGAQQLQEQLPLIVGSATAGLVFVVAVVVIAIV ECGPCEASVRYSEPPHGLTRTSVTVSDLEPHMNYTFTVEARNGVSGLVTSRSFR.TASVS..I..NQ...TEPPKVRLEGRSTTSLSVSW.SIPPPQQSR il. YEvk $\gamma\gamma$ ekdq. ersy. iv..k. tsvt. dgLkpdt. YvfqvrarTaaGyG.. Sr.. efeT. pea. sgsg \dots ivvviivs. aga \dots ll $vv\dots$ v.l \dots r MKYEV. TYRKKGDSNSYNVRRTEGFSVTLDDLAPDTTYLVQVQALTQEGQGAGSKVHEFQTLSPEGSGNLAVIGGVAVGVVLLLVLAGVGFFIHRRKKN PCQPCGVGVHFSPGARALTTPAVHVNGLEPYANYTFNVEAQNGVSGLGSSGHAS..TSVSISMGHAESLS..GLSLRLVKKEPRQLELTWAGSRPRSPGA ECVPCGSNIGYMPQQTGLEDNYVTVMDLLAHANYTFEVEAVNGVSDL....SRSQRLFAAVSITTGQAAPSQVSGVMKERVLQRSVELSW.QEPEHPNGV QCEPCSPNVRFLPRQFGLTNTTVTVTDLLAHTNYTFEIDAVNGVSEL..SSPPRQFAAV..SITTNQAAPSPVLTIKKDRTSRNSISLSW.QEPEHPNGI ACTRCGDNVQYAPRQLGLTEPRIYISDLLAHTQYTFEIQAVNGVTD..QSPFSPQFASV..NITTNQAAPSAVSIMHQVSRTVDSITLSW.SQPDQPNGV /CEECGGHVRYLPRQSGLKNTSVMMVDLLAHTNYTFEIEAVNGVSDL....SPGARQYVSVNVTTNQAAPSPVTNVKKGKIAKNSISLSM.QEPDRPNGI KCRPCGSGVHYTPQQÑGLKTTKVSITDLLAHTNYTFEIWAVNGVSK....YNPNPDQSVSVTVTTNQAAPSSIALVQAKEVTRYSVALAW.LEPDRPNGV acsrcddnvefvprolglseprvhtshllahtrytfevoavngvsgk....splppryaavnittnqaapsevptlrlhsssgssltlsw.apperpngv CepCg.nvry.prq1gLt.t.vtvsdLlahtnYtFe.eAvNGVs.l....sp.q.asvsv.ittnqaaps.v.tvr....sr.s.s1sW.qep.rpngv HEX11 CONS HEK4 HEK5 HEK7 HEK8 HEK2 HEK4 HEK5 HEK7 JEK8 HEK2 CONS EPH ECK SUBSTITUTE SHEET (RULE 26)

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?RQRQQRHVTAPPMWIERTSCAEALCGTSRHTRTLHREPWTL..PGGWSNFPSRELDPAWLMVDTVIGEGEFGEVYRGTLRLPS.QDCKTVAIKTLKDTS PGGQWWNFLREATIMGQFSHPHILHLEGVVTKRKPIMIITEFMENAALDAFLREREDQLVPGQLVAMLQGIASGNNYLSNHNYVHRDLAARNILVNQNLC :EKQRVDFLGEAGIMGQFSHHNIIRLEGVISKYKPMMIITEYMENGALDKFLREKDGEFSVLQLVGMLRGIAAGMKYLANMNYVHRDLAARNILVNSNLV EKQRRDFLGEASIMGQFDHPNIIRLEGVVTKSKPVMIVTEYMENGSLDSFLRKHDAQFTVIQLVGMLRGIASGMKYLSDMGYVHRDLAARNILINSNLV: $\tt EKQRRDFLSEASIMGQFDHPNVIHLEGVVTKSTPVMIITEFMENGSLDSFLRQNDGQFTVIQLVGMLRGIAAGMKYLADMNYVHRDLAARNILVNSNLV$ FEKQRRDFLGEASIMGQFDHPNIIHLEGVVTKSKPVMIVTEYMENGSLDTFLKKNDGQFTVIQLVGMLRGISAGMKYLSDMGYVHRDLAARNILINSNLV IDKQRRDFLSEASIMGQFDHPNIIHLEGVVTKCKPVMIITEYMENGSLDAFLRKNDGRFTVIQLVGMLRGIGSGMKYLSDMSYVHRDLAARNILVNSNLV 'ERQRRDFLSEASIMGQFDHPNIIRLEGVVTKSRPVMILTEFMENCALDSFLRLNDGQFTVIQLVGMLRGIAAGMKYLSEMNYVHRDLAARNILVNSNLV TEKQRRDFLCEASIMGQFDHPNVVHLEGVVTRGKPVMIVIEFMENGALHAFLRKHDGQFTVIQLVGMLRGIAAGMRYLADMGYVHRDLAARNILVNSNLV r..gsr.dd.ey.keq.....klpg.ktyidP.TyedPngav.efakEidascikiekviGaGEFGEVcsGrLklp.gkre..VAIKTLKvgyLKPLKTYVDPHTYEDPNQAVLKFTTEIHPSCVTRQKVIGAGEFGEVYKGMLKTSSGKKEVPVAIKTLKAGY °CGYKSKHGADEKRLHFGNG.....HLKLPGLRTYVDPHTYEDPTQAVHEFAKELDATNISIDKVVGAGEFGEVCSGRLKLPS.KKEISVAIKTLKVGY NRRGFERADSEYTDKLQHYT.....SGHITPGMKIYIDPFTYEDPNEAVREFAKEIDISCVKIEQVIGAGEFGEVCSGHLKLP.GKREIFVAIKTLKSGY RCGYSKAKQDPEEEKMHFHN.....GHIKLPGVRTYIDPHTYEDPNQAVHEFAKEIEASCITIERVIGAGEFGEVCSGRLKLP.GKRELPVAIKTLKVGY RRRSKYSKAKQEADEEKHLN............QGVRTYVDPFTYEDPNQAVREFAKEIDASCIKIEKVIGVGEFGEVCSGRLKVP.GKREICVAIKTLKAGY CLRKQRHGSDSEYTEKLQQY......IAPGMKVYIDPFTYEDPNEAVREFAKEIDVSCVKIEEVIGAGEFGEVCRGRLKQP.GRREVFVAIKTLKVGY VFGFIIGRRHCGYTKADQEGDEELYFHFKFPGTKTYIDPETYEDPNRAVHQFAKELDASCIKIERVIGAGEFGEVCSGRLKLP.GKRDVAVAIKTLKVGY cekQrrdFL.EasIMGQFdHpniihLEGVvtkskPvMIitE.MENg.Ld.FLrkndgqftviQLVgMLrGIaaGMkYLsdmnYVHRDLAARNILvNsNLv DRARQSPEDVYFSKSEQ. HEK11 HEK4 HEK5 HEX8 HEK7 HEK2 HEK4 **JEKS** TEK7 1EK8 IEK2 ECK EPH ECK SUBSTITUTE SHEET (RULE 26)

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CKVSDFG1sRv1eDD.pea.yT.trGGkiPiRwTaPEAIayRkFTsASDVWSyGIVmWEVmsyGerPYw.msNqdVikaieegyRLPpPmDCPaal.qLM CKVSDFGLTRLL.DDFDGTYET..QGGKIPIRWTAPEAIAHRIFTTASDVWSFGIVMWEVLSFGDKPYGEMSNQEVMKSIEDGYRLPPPVDCPAPLYELM :KVSDFGLSRVLEDD. PEATYT. TSGGKIPIRWTAPEAISYRKFTSASDVWSFGIVMWEVMTYGERPYWELSNHEVMKAINDGFRLPTPMDCPSAIYQLM CKVSDFGLSRVLEDD. PEAAYT. TRGGKIPIRWTSPEAIAYRKFTSASDVWSYGIVLWEVMSYGERPYWEMSNQDVIKAVDEGYRLPPPMDCPAALYQLM :KVSDFGLSRFLEDDTSDPTYTSALGGKFPIRWTAPEAIQYRKFTSASDVWSYGIVMWEVMSYGERPYWDMTNQDVINAIEQDYRLPPPMDCPSALHQLM JKVSDFGLSRVLEDD. PEAAYT. TRGGKIPIRWTAPEAIAFRKFTSASDVWSYGIVMWEVVSYGERPYWEMTNQDVIKAVEEGYRLPSPMDCPAALYQLM CKVSDFGMSRVLEDD. PEAAYT. TRGGKI PIRWTA PEAIA YRKFTSA SDVWSYGIVMWEVMSYGER PYWDMSNQDVIKA I EEGYRL PPPMDCPIALHQLM CKVSDFGLSRFLEDDPSDPTYTSSLGGKIPIRWTAPEAIAYRKFTSASDVWSYGIVMWEVMSYGERPYWDMSNQDVINAVEQDYRLPPPMDCPTALHQLM CKVSDFGLSRVIEDD. PEAVYT. TTGGKIPVRWTAPEAIQYRKFTSASDVWSYGIVMWEVMSYGERPYWDMSNQDVIKAIEEGYRLPAPMDCPAGLHQLM HEK11 CONS HEK4 HEK5 HEK7 HEK8 HEK2 EPH

IQCWQQERARRPKFADIVSILDKLIRAPDSLKTLADFDPRVSIRLPSTSGSEGVPFRTVSEWLESIKMQQYTEHFWAAGYTAIEKVVQMTNDDIKRIGVR JDCWQKERNSRPKFDEIVNMLDKLIRNPSSLKTLVNASCRVSNLLAEHSPLGSGAYRSVGEWLEAIKMGRYTEIFMENGYSSMDAVAQVTLEDLRRLGVT JDCWVRDRNLRPKFSQIVNTLDKLIRNAASLKVIASAQSGMSQPLLDRTVPDYTTFTTVGDWLDAIKMGRYKESFVSAGFASFDLVAQMTAEDLLRIGVT LDCWQKERAERPKFEQIVGILDKMIRNPNSLKTPLGTCSRPISPLLDQNTPDFTTFCSVGEWLQAIKMERYKDNFTAAGYNSLESVARMTIEDVMSLGIT .dCWqk.RnrRPkF.qivniLdklirnpnSLktia.assr.s.pLld.sgpd.ttfrtvgeWLeaikmgryke.Ftaagyts..avaqmtaeDl.riGvt ONCWAYDRARRPHFQKLQAHLEQLLANPHSLRTIANFDPRVTLRLPSLSGSDGIPYRTVSEWLESIRMKRYILHFHSAGLDTMECVLELTAEDLTQMGIT JOCWQKDRINNRPKFEQIVSILDKLIRNPGSLKIITSAAARPSNLLLDQSNVDISTFRTTGDWLNGVRTAHCKEIFTGVEYSSCDTIAKISTDDMKKVGVT JDCWQKDRNHRPKFGQIVNTLDKMIRNPNSLKAMAPLSSGINLPLLDRTIPDYTSFNTVDEWLEAIKMGQYKESFANAGFTSFDVVSQMMMEDILRVGVT JDCWQKERSDRPKFGQIVNMLDKLIRNPNSLKRTGTESSRPNTALLDPSSPEFSAVVSVGDWLQAIKMDRYKDNFTAAGYTTLEAVVHVNQEDLARIGIT CONS HEK4 **JEKS** 1EK8 **JEK2** 1EK7 ECK

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F1G. 5F

CONS lvghQkilsSiq.mr.Qmnqgh.p.v.V
EPH LPGHQKRILCSIQGFKD
ECK LPGHQKRIAYSLLGLKDQVNTVGIPI
HEK4 VVGPQKKIISSIKALETQSKNGPVPV
HEK5 LAGHQKKILNSIQVMRAQMNQIQSVEV
HEK7 LVGHQKKIMNSLQEMKVQLVNGMVPL
HEK8 AITHQNKILSSVQAMRTQMQQMHGRMVPV
HEK2 LAGHQKKILSSIQDMRLQMNQTLPVQV
HEK11 LVGHQKKIMSSIQTMRAQMLHLHGTGIQV

28/3**3** FIG. 6

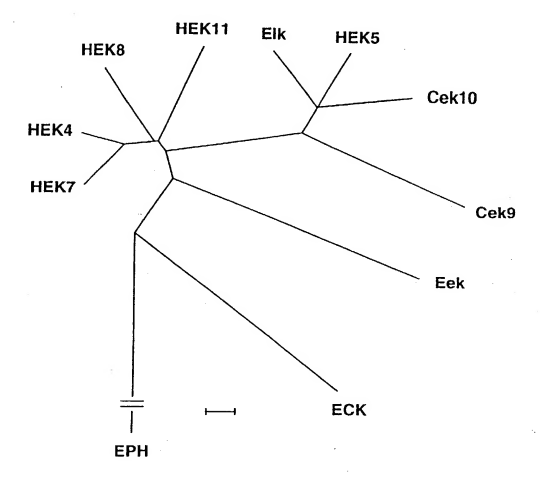


FIG. 7A

<u>Human</u>

Hear air certa

FIG. 7B

Rat

9.5 kb — 7.5 — 1.5

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FIG. 8A

<u>Human</u>

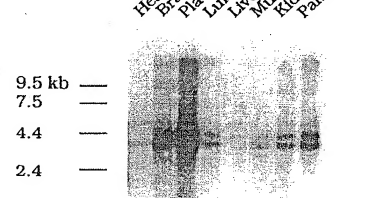


FIG. 8B

Rat

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FIG. 9A

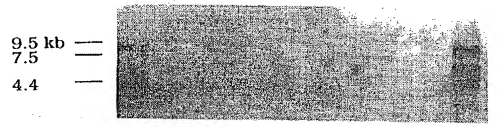
<u>Human</u>

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FIG. 9B

Rat

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FIG. IOA

<u>Human</u>

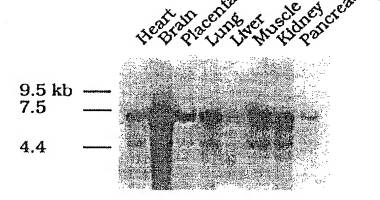


FIG. IOB

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Ovary estis trythus theat stornacht ture lines tidney air



FIG. 11A

<u>Human</u>

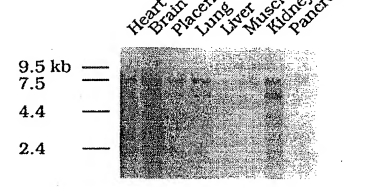
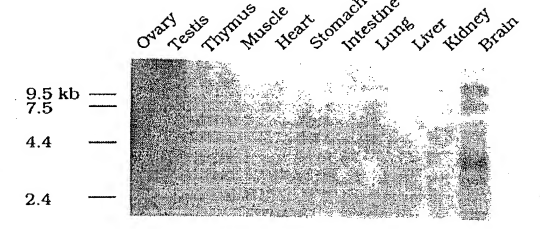


FIG. IIB

<u>Rat</u>



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INTERNATIONAL SEARCH REPORT

Interr al Application No PCT/US 95/04681

A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 C12N15/12 C07K14/71 C07K16/28 A61K39/395 A61K38/17 C12N15/62 G01N33/566 According to International Patent Classification (IPC) or to both national classification and IPC Minimum documentation searched (classification system followed by classification symbols) C12N C07K A61K G01N IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X WO-A-93 00425 (INST MEDICAL W & E HALL) 7 1-8,10, 15-18, January 1993 20,23, 25-32,34 see the whole document X DE-A-42 33 782 (CHEMOTHERAPEUTISCHES 1-9, 15-19, FORSCHUNG) 14 April 1994 23, 25-32,34 see the whole document X CA-A-2 083 521 (MOUNT SINAI HOSPITAL CORP 1-7,13, 15-18,) 1 October 1993 23-32,34 see the whole document -/--Further documents are listed in the continuation of box C. Patent family members are listed in annex. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance invention .E. earlier document but published on or after the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. document referring to an oral disclosure, use, exhibition or document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 15. 09. 95 6 September 1995 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Nauche, S

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Interr 1al Application No
PCT/US 95/04681

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Interr nal Application No
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C (Continue	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	PC1/03 35/04001
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
х	ONCOGENE, vol. 6, no. 6, 1991 pages 1057-1061, CHAN, J.; WATT, V.M.; 'eek and erk, new members of the eph subclass of receptor protein-tyrosine kinases' cited in the application see the whole document	1-9, 15-18, 23, 25-27, 32,34
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ernational application No.

INTERNATIONAL SEARCH REPORT

PCT/US 95/04681

Box I Obse	ervations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
This internation	onal search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
Rem.	use they relate to subject matter not required to be searched by this Authority, namely: ark: Although claim 32 is directed to a method of treatment of the human/animal body (Rule 39.1(iv)) PCT), the search has been carried out and based on the alleged effects of the compound/composition. as Nos: use they relate to parts of the international application that do not comply with the prescribed requirements to such tent that no meaningful international search can be carried out, specifically:	
becau	ns Nos.: use they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a). Transverse unity of invention is lacking (Continuation of item 2 of first sheet)	
This Internation	onal Searching Authority found multiple inventions in this international application, as follows:	$\frac{1}{1}$
1. As all search	required additional search fees were timely paid by the applicant, this international search report covers all able claims.	
2. As all of any	searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment additional fee.	
3. As onl covers	ly some of the required additional search fees were timely paid by the applicant, this international search report only those claims for which fees were paid, specifically claims Nos.:	
4. No req	quired additional search fees were timely paid by the applicant. Consequently, this international search report is ed to the invention first mentioned in the claims; it is covered by claims Nos.:	
Remark on Prote	The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.	

INTERNATIONAL SEARCH REPORT

information on patent family members

Inten nal Application No
PCT/US 95/04681

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
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DE-A-4233782	14-04-94	NONE		
CA-A-2083521		NONE		

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